PCT PCT

INTERNATIONAL SEARCH REPORT

(PCT Article 18 and Rules 43 and 44)

Applicant's or agent's file reference P23667A/RMC	FOR FURTHER see Notification of Transmittal of International Search Report (Form PCT/ISA/220) as well as, where applicable, item 5 below.						
International application No.	International filing date (day/month/year)	(Earliest) Priority Date (day/month/year)					
PCT/GB 00/01089	29/03/2000 29/03/1999						
Applicant UNIVERSITY OF ULSTER et a	1.						
according to Article 18. A copy is being to This International Search Report consists	ansmitted to the International Bureau.	Authority and is transmitted to the applicant this report.					
Basis of the report a. With regard to the language, the	international search was carried out on the	basis of the international application in the					
the international search wathority (Rule 23.1(b)). b. With regard to any nucleotide as was carried out on the basis of the contained in the international filed together with the international subsequently to the statement that the substantian international application of the statement that the informational application is the statement that the information is the statement that the	nd/or amino acid sequence disclosed in the sequence listing: conal application in written form. ernational application in computer readable to this Authority in written form. to this Authority in computer readble form. becausely furnished written sequence listings filed has been furnished. ormation recorded in computer readable formation recorded in computer readable formation unsearchable (See Box I).	of the international application furnished to this ne international application, the international search form. In does not go beyond the disclosure in the rm is identical to the written sequence listing has been					
the text has been establi	ubmitted by the applicant. shed by this Authority to read as follows: HIBITORY PEPTIDE AND THEI	R USE FOR TREATMENT OF DIABETES					
the text has been establi	ubmitted by the applicant. shed, according to Rule 38.2(b), by this Au e date of mailing of this international searc	thority as it appears in Box III. The applicant may, n report, submit comments to this Authority.					
as suggested by the app because the applicant fa		None of the figures.					

INTERNATIONAL SEARCH REPORT

nternatio	nal	Application No
P	В	00/01089

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C07K14/575 A61K38/22 A61P5/48

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) IPC 7 C07K A61K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

CHEM ABS Data, EPO-Internal, WPI Data, PAJ, BIOSIS

ategory °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
1	DE 196 16 486 A (HANS KNOELL INST FUER NATURSTO) 30 October 1997 (1997-10-30) claims; examples	1,7-11
4	EP 0 869 135 A (LILLY CO ELI) 7 October 1998 (1998-10-07) page 3, line 2 - line 49; claims; examples	1,7-11
		
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X Further documents are listed in the continuation of box C.	Patent family members are listed in annex.
"Special categories of cited documents: "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or	 "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
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Date of the actual completion of the international search 1 November 2000	Date of mailing of the international search report 15/11/2000
Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL – 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authorized officer Fuhr, C



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INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)

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opplicant's or agent's file reference	FOR FURTHER ACTION	See Notifica Preliminary	Examination	1
P23667A/CPA/RMC	International filing date (day/month	h/year)	Priority date (day/month/year) 29/03/1999	
International application No.	29/03/2000		29/00/100	. 1
PCT/GB00/01089 International Patent Classification (IPC) o	r national classification and IPC			13
International Patent Classification (5 C) C07K14/575				
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This international preliminary of and is transmitted to the application.	examination report has cant according to Article 36.		nternational Preliminary Examining Authority	
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3. This report contains indica	tions relating to the following items			
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Basis of the re	eport ment of opinion with regard to nover invention	alty inventiv	ve step and industrial applicability	
II Priority	nment of opinion with regard to nov	eity, inves	industrial applicability;	
III Norrestablic	of invention	oard to nove	elty, inventive step or industrial application	
V Reasoned st	atement under Article 35(2) with re lexplanations suporting such state	ment	elty, inventive step or industrial applicability;	
Citations	hotiod			
VI Certain defe	uments cited acts in the international application	ication		
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Fax: +49 89 23	99 - 4700			
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INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No. PCT/GB00/01089

I. Basis	of the	report
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ı.	Basis	s of the report	the boundary from the section of the
1.	the re	The Office in	nents of the international application (Replacement sheets which have been furnished to response to an invitation under Article 14 are referred to in this report as "originally filed" this report since they do not contain amendments (Rules 70.16 and 70.17)):
	1-32		as originally filed
	Clain	ns, No.:	
	1-11		as originally filed
	Draw	vings, sheets:	
	1/32-	-32/32	as originally filed
	Sequ	uence listing par	t of the description, pages:
	1,2,1	filed with the letter	r of 06.07.00
2.	With lang	regard to the lan uage in which the	guage, all the elements marked above were available or furnished to this Authority in the international application was filed, unless otherwise indicated under this item.
	Thes	se elements were	available or furnished to this Authority in the following language: , which is:
		the language of a	translation furnished for the purposes of the international search (under Rule 23.1(b)).
	П	the language of D	publication of the international application (under Rule 48.3(b)).
		the language of a 55.2 and/or 55.3)	a translation furnished for the purposes of international preliminary examination (under Rule
3	. With inter	n regard to any nu rnational prelimina	icleotide and/or amino acid sequence disclosed in the international application, the ary examination was carried out on the basis of the sequence listing:
		contained in the i	international application in written form.
		filed together with	h the international application in computer readable form.
	\boxtimes	furnished subsec	quently to this Authority in written form.
	\boxtimes	furnished subsec	quently to this Authority in computer readable form.
	×	The statement the	nat the subsequently furnished written sequence listing does not go beyond the disclosure in application as filed has been furnished.
	×	The statement the	nat the information recorded in computer readable form is identical to the written sequence

4. The amendments have resulted in the cancellation of:

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INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No. PCT/GB00/01089

		the description,	pages:								
		the claims,	Nos.:								
		the drawings,	sheets:								
5.		This report has been considered to go bey						ad not be	en made,	, since the	y have beer
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6.	Add	itional observations, it	necessar	y:							
V.		soned statement un tions and explanatio					elty, inve	entive ste	p or indu	ıstrial apı	plicability;
1.	Stat	ement									
	Nov	elty (N)	Yes: No:	Claims Claims	1-11						

2. Citations and explanations see separate sheet

Industrial applicability (IA)

Inventive step (IS)

Yes:

No:

Yes:

No:

Claims 1-11

Claims 1-11 Claims

Claims

D1: DE-A-196 16 486

The known Gastric Inhibitory Polypeptide (GIP) contains 1 to 42 amino acid residues (GIP (1-42)) and acts as an insulin-releasing hormone. This hormone can be inactivated by enzymatic degradation by a dipeptidyl peptidase-IV (DPP-IV). The present invention provides peptide analogues of GIP which are DPP-IV resistant and can therefore be used in the treatment of diabetes.

The subject-matter of <u>claims 1-11</u> which is directed to said peptide analogues of GIP, to the use thereof, pharmaceutical compositions and methods, appears to be novel and to involve an inventive step (Article 33(2)(3) PCT) since the prior art does not disclose peptide GIPs modified in particular in the N-terminal amino acid residues at positions 1-3 as presently claimed. Moreover, as also indicated by the Applicant, it was not established in 1999 that DPPIV is a major inactivator of GIP in vivo, nor was it established that the N-terminal amino acid portion of GIP is a key for hormone activity.

The above comments are based on the assumption that all claims enjoy priority rights from the filing date of the priority document. In that case the document cited as P,X-document in the international search report is not considered as prior art.

The prior art document D1 is not mentioned in the description, cf. Rule 5.1(a)(ii) PCT. Moreover, a document reflecting the prior art described on pages 1-4 is not identified in the description (Rule 5.1(a)(ii) PCT).

INTERNATION EARCH REPORT



nal Application No

PCT/GB 00/01089 A. CLASSIFICATION OF SUBJECT MATTER IPC 7 CO7K14/575 A61K38/22 A61P5/48 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) C07K A61K IPC 7 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) CHEM ABS Data, EPO-Internal, WPI Data, PAJ, BIOSIS C. DOCUMENTS CONSIDERED TO BE RELEVANT Category 9 Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. A DE 196 16 486 A (HANS KNOELL INST FUER 1,7-11-NATURSTO) 30 October 1997 (1997-10-30) claims; examples A EP 0 869 135 A (LILLY CO ELI) 1,7-117 October 1998 (1998-10-07) page 3, line 2 - line 49; claims; examples -/--Further documents are listed in the continuation of box C. Х X Patent family members are listed in annex. Special categories of cited documents : T' later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the "O" document referring to an oral disclosure, use, exhibition or document is combined with one or more other such docu-ments, such combination being obvious to a person skilled in the art. document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 1 November 2000 15/11/2000 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl,

Fax: (+31-70) 340-3016

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Interne al Application No PCT/GB 00/01089

C.(Continue	C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT			
Category *	Citation of document, with indication, where appropriate, of the relevant passages		Relevant to claim No.	
Ρ,Χ	CHEMICAL ABSTRACTS, vol. 131, no. 1, 5 July 1999 (1999-07-05) Columbus, Ohio, US; abstract no. 944, 0'HARTE, FINBARR P. M. ET AL: "NH2-terminally modified gastric inhibitory polypeptide exhibits aminopeptidase resistance and enhanced antihyperglycemic activity" XP002151675 abstract & DIABETES (4-1999), 48(4), 758-765, 1999,		1,7-11	
PCT/ISA/210 (o	ontinustion of second sheets (July 1992)			



...ormation on patent family members

Internation No. PCT/GB 00/01089

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Patent document cited in search report		Publication date		atent family member(s)	Publication date
DE 19616486	A	30-10-1997	AU AU CA CN WO EP NZ	721477 B 3023397 A 2252576 A 1216468 A 9740832 A 0896538 A 332707 A	06-07-2000 19-11-1997 06-11-1997 12-05-1999 06-11-1997 17-02-1999 28-10-1999
EP 0869135	Α	07-10-1998	US	5981488 A	09-11-1999





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(54) Title: PEPTIDE

(57) Abstract

The present invention provides peptides which stimulate the release of insulin. The peptides, based on GIP 1-42 include substitutions and/or modifications which enhance and influence secretion and/or have enhanced resistance to degradation. The invention also provides a process of N terminally modifying GIP and the use of the peptide analogues for treatment of diabetes.

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1

"Peptide"

1

2 The present invention relates to the release of insulin 3 and the control of blood glucose concentration. More 4 particularly the invention relates to the use of 5 peptides to stimulate release of insulin, lowering of 6 blood glucose and pharmaceutical preparations for 7 treatment of type 2 diabetes. 8 9 Gastric inhibitory polypeptide (GIP) and glucagon-like 10 peptide-1(7-36)amide (truncated GLP-1; tGLP-1) are two 11 important insulin-releasing hormones secreted from 12 endocrine cells in the intestinal tract in response to 13 Together with autonomic nerves they play a 14 vital supporting role to the pancreatic islets in the 15 control of blood glucose homeostasis and nutrient 16 17 metabolism. 18 Dipeptidyl peptidase IV (DPP IV; EC 3.4.14.5) has been 19 identified as a key enzyme responsible for inactivation 20 of GIP and tGLP-1 in serum. DPP IV is completely 21 inhibited in serum by the addition of diprotin A(DPA, 22 0.1 mmol/l). This occurs through the rapid removal of 23

2

the N-terminal dipeptides Tyr1- Ala2 and His7-Ala8 1 giving rise to the main metabolites GIP(3-42) and GLP-2 These truncated peptides 1(9-36)amide, respectively. 3 are reported to lack biological activity or to even 4 serve as antagonists at GIP or tGLP-1 receptors. The 5 resulting biological half-lives of these incretin 6 hormones in vivo are therefore very short, estimated to 7 be no longer than 5 min. 8 9 In situations of normal glucose regulation and 10 ' pancreatic B-cell sensitivity, this short duration of 11 action is advantageous in facilitating momentary 12 adjustments to homeostatic control. However, the 13 current goal of a possible therapeutic role of incretin 14 hormones, particularly tGLP-1 in NIDDM therapy is 15 frustrated by a number of factors in addition to 16 finding a convenient route of administration. Most 17 notable of these are rapid peptide degradation and 18 rapid absorption (peak concentrations reached 20 min) 19 and the resulting need for both high dosage and precise 20 timing with meals. Recent therapeutic strategies have 21 focused on precipitated preparations to delay peptide 22 absorption and inhibition of GLP-1 degradation using 23 specific inhibitors of DPP IV. A possible therapeutic 24 role is also suggested by the observation that a 25 specific inhibitor of DPP IV, isoleucine thiazolidide, 26 lowered blood glucose and enhanced insulin secretion in 27 glucose-treated diabetic obese Zucker rats presumably 28 by protecting against catabolism of the incretin 29 hormones tGLP-1 and GIP. 30 31

3

.

Numerous studies have indicated that tGLP-1 infusion 1 restores pancreatic B-cell sensitivity, insulin 2 secretory oscillations and improved glycemic control in 3 various groups of patients with IGT or NIDDM. Longer 4 term studies also show significant benefits of tGLP-1 5 injections in NIDDM and possibly IDDM therapy, 6 providing a major incentive to develop an orally 7 effective or long-acting tGLP-1 analogue. 8 attempts have been made to produce structurally 9 modified analogues of tGLP-1 which are resistant to DPP 10 IV degradation. A significant extension of serum half-11 life is observed with His7- glucitol tGLP-1 and tGLP-1 12 analogues substituted at position 8 with Gly, Aib, Ser 13 However, these structural modifications seem 14 to impair receptor binding and insulinotrophic activity 15 thereby compromising part of the benefits of protection 16 from proteolytic degradation. In recent studies using 17 His⁷-glucitol tGLP-1, resistance to DPP IV and serum 18 degradation was accompanied by severe loss of insulin-19 20 releasing activity. 21 GIP shares not only the same degradation pathway as 22 tGLP-1 but many similar physiological actions, 23 including stimulation of insulin and somatostatin 24 secretion, and the enhancement of glucose disposal. 25 These actions are viewed as key aspects in the 26 antihyperglycemic properties of tGLP-1, and there is 27 therefore good expectation that GIP may have similar 28 potential as NIDDM therapy. Indeed, compensation by 29 GIP is held to explain the modest disturbances of 30 glucose homeostasis observed in tGLP-1 knockout mice. 31 Apart from early studies, the anti-diabetic potential 32

4

of GIP has not been explored and tGLP-1 may seem more 1 2 attractive since it is viewed by some as a more potent 3 insulin secretagogue when infused at "so called" physiological concentrations estimated by RIA. 4 5 6 The present invention aims to provide effective analogues of GIP. It is one aim of the invention to 7 provide a pharmaceutical for treatment of Type 2 8 9 diabetes. 10 According to the present invention there is provided an 11 effective peptide analogue of the biologically active 12 GIP(1-42) which has improved characteristics for 13 treatment of Type 2 diabetes wherein the analogue 14 comprises at least 15 amino acid residues from the N 15 terminus of GIP(1-42) and has at least one amino acid 16 substitution or modification at position 1-3 and not 17 18 including Tyr1 glucitol GIP(1-42). 19 The structures of human and porcine GIP(1-42) are shown 20 The porcine peptide differs by just two amino 21 acid substitutions at positions 18 and 34. 22 23 24 25 The analogue may include modification by fatty acid 26 addition at an epsilon amino group of at least one lysine residue. 27 28 29 The invention includes Tyr¹ glucitol GIP(1-42) having 30 fatty acid addition at an epsilon amino group of at 31 least one lysine residue. 32

5

Fig. 1. Primary structure of human gastric inhibitory polypeptide (GIP)

1 5 10 15 20 25 NH₂-Tyr-Ala-Glu-Gly-Thr-Phe-Ile-Ser-Asp-Tyr-Ser-Ile-Ala-Met-Asp-Lys-Ile-His-Gln-Gln-Asp-Phe-Val-Asn-Trp-

Leu-Leu-Ala-Gln-Lys-Gly-Lys-Lys-Asn-Asp-Trp-Lys-His-Asn-Ile-Thr-Gln-COOH

Fig. 2. Primary structure of porcine gastric inhibitory polypeptide (GIP)

1 5 10 15 20 25 NH₂-Tyr-Ala-Glu-Gly-Thr-Phe-Ile-Ser-Asp-Tyr-Ser-Ile-Ala-Met-Asp-Lys-Ile-<u>Arg</u>-Gln-Gln-Asp-Phe-Val-Asn-Trp-

Lou-Leu-Ala-Gln-Lys-Gly-Lys-Lys-Ser-Asp-Trp-Lys-His-Asn-Ile-Thr-Gln-COOH

- 1 Analogues of GIP(1-42) may have an enhanced capacity to
- 2 stimulate insulin secretion, enhance glucose disposal,
- 3 delay glucose absorption or may exhibit enhanced
- 4 stability in plasma as compared to native GIP. They
- 5 also may have enhanced resistance to degradation.

6

- 7 Any of these properties will enhance the potency of the
- 8 analogue as a therapeutic agent.

9

- 10 Analogues having D-amino acid substitutions in the 1, 2
- 11 and 3 positions and/or N-qlycated, N-alkylated, N-
- 12 acetylated or N-acylated amino acids in the 1 position
- 13 are resistant to degradation in vivo.

14

- 15 Various amino acid substitutions at second and third
- 16 amino terminal residues are included, such as GIP(1-
- 17 42)Gly2, GIP(1-42)Ser2, GIP(1-42)Abu2, GIP(1-42)Aib,
- 18 GIP(1-42)D-Ala2, GIP(1-42)Sar2, and GIP(1-42)Pro3.

- 20 Amino-terminally modified GIP analogues include N-
- 21 glycated GIP(1-42), N-alkylated GIP(1-42), N-actylated

```
GIP(1-42), N-acetyl-GIP(1-42) and N-isopropyl GIP(1-
 1
 2
     42).
 3
    Other stabilised analogues include those with a peptide
 4
     isostere bond between amino terminal residues at
 5
    position 2 and 3. These analogues may be resistant to
 6
     the plasma enzyme dipeptidyl-peptidase IV (DPP IV)
 7
     which is largely responsible for inactivation of GIP by
 8
     removal of the amino-terminal dipeptide Tyrl-Ala2.
 9
10
     In particular embodiments, the invention provides a
11
    peptide which is more potent than human or porcine GIP
12
     in moderating blood glucose excursions, said peptide
13
     consisting of GIP(1-42) or N-terminal fragments of
14
    GIP(1-42) consisting of up to between 15 to 30 amino
15
     acid residues from the N-terminus (i.e. GIP(1-15) -
16
    GIP(1-3)) with one or more modifications selected from
17
18
     the group consisting of:
19
          substitution of Ala<sup>2</sup> by Gly
20
     (a)
          substitution of Ala<sup>2</sup> by Ser
21
     (b)
          substitution of Ala<sup>2</sup> by Abu
     (c)
22
          substitution of Ala<sup>2</sup> by Aib
     (d)
23
          substitution of Ala2 by D-Ala
24
     (e)
          substitution of Ala<sup>2</sup> by Sar
     (f)
25
          substitution of Glu<sup>3</sup> by Pro
26
     (g)
          modification of Tyr1 by acetylation
27
     (h)
28
     (i)
          modification of Tyr1 by acylation
          modification of Tyr1 by alkylation
     (j)
29
          modification of Tyr1 by glycation
     (k)
30
          conversion of Ala<sup>2</sup>-Glu<sup>3</sup> bond to a psi [CH2NH] bond
31
     (1)
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conversion of Ala2-Glu3 bond to a stable peptide (m) 1 isotere bond 2 (n) (n-isopropyl-H) 1GIP. 3 4 The invention also provides the use of Tyr1-glucitol 5 GIP in the preparation of a medicament for the 6 treatment of diabetes. 7 8 The invention further provides improved pharmaceutical 9 compositions including analogues of GIP with improved 10 pharmacological properties. 11 12 Other possible analogues include certain commonly 13 encountered amino acids, which are not encoded by the 14 genetic code, for example, beta-alanine (beta-ala), or 15 other omega-amino acids, such as 3-amino propionic, 4-16 amino butyric and so forth, ornithine (Orn), citrulline 17 (Cit), homoarginine (Har), t-butylalanine (t-BuA), t-18 butylglycine (t-BuG), N-methylisoleucine (N-MeIle), 19 phenylglycine (Phg), and cyclohexylalanine (Cha), 20 norleucine (Nle), cysteic acid (Cya) and methionine 21 sulfoxide (MSO), substitution of the D form of a 22 neutral or acidic amino acid or the D form of tyrosine 23 for tyrosine. 24 25 According to the present invention there is also 26 provided a pharmaceutical composition useful in the 27 treatment of diabetes type II which comprises an 28 effective amount of the peptide as described herein, in 29 admixture with a pharmaceutically acceptable excipient. 30

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8 The invention also provides a method of N-terminally 1 modifying GIP or analogues thereof the method 2 comprising the steps of synthesizing the peptide from 3 the C terminal to the penultimate N terminal amino 4 acid, adding tyrosine to a bubbler system as a F-moc 5 protected Tyr(tBu)-Wang resin, deprotecting the N-6 terminus of the tyrosine and reacting with the 7 modifying agent, allowing the reaction to proceed to 8 completion, cleaving the modified tyrosine from the 9 Wang resin and adding the modified tyrosine to the 10 peptide synthesis reaction. 11 12 Preferably the agent is glucose, acetic anhydride or 13 pyroglutamic acid. 14 15 The invention will now be demonstrated with reference 16 to the following non-limiting example and the 17 accompanying figures wherein: 18 19 Figure 1a illustrates degradation of GIP by DPP IV. 20 21 Figure 1b illustrates degradation of GIP and Tyr1 22 glucitol GIP by DPP IV. 23 24 Figure 2a illustrates degradation of GIP human plasma. 25 26

Figure 2b illustrates degradation of GIP and Tyr1-

glucitol GIP by human plasma.

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Figure 3 illustrates electrospray ionization mass
    spectrometry of GIP, Tyr1-glucitol GIP and the major
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    degradation fragment GIP(3-42).
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    Figure 4 shows the effects of GIP and glycated GIP on
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    plasma glucose homeostasis.
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    Figure 5 shows effects of GIP on plasma insulin
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    responses.
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    Figure 6 illustrates DPP-IV degradation of GIP 1-42.
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    Figure 7 illustrates DPP-IV degradation of GIP (Abu^2).
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    Figure 8 illustrates DPP-IV degradation of GIP (Sar^2).
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    Figure 9 illustrates DPP-IV degradation of GIP (Ser2),
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    Figure 10 illustrates DPP-IV degradation of N-Acetyl-
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20
    GIP.
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    Figure 11 illustrates DPP-IV degradation of glycated
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23
    GIP.
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    Figure 12 illustrates human plasma degradation of GIP.
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    Figure 13 illustrates human plasma degradation of GIP
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    (Abu<sup>2</sup>).
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    Figure 14 illustrates human plasma degradation of GIP
30
    (Sar^2).
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Figure 15 illustrates human plasma degradation of GIP 1 2 (Ser^2) . 3 4 Figure 16 illustrates human plasma degradation of glycated GIP. 5 6 Figure 17 shows the effects of various concentrations 7 of GIP 1-42 and GIP (Abu²) on insulin release from 8 BRIN-BD11 cells incubated at 5.6mM glucose. 9 10 Figure 18 shows the effects of various concentrations 11 of GIP 1-42 and GIP (Abu²) on insulin release from 12 BRIN-BD11 cells incubated at 16.7mM glucose. 13 14 Figure 19 shows the effects of various concentrations 15 of GIP 1-42 and GIP (Sar2) on insulin release from 16 BRIN-BD11 cells incubated at 5.6mM glucose. 17 18 Figure 20 shows the effects of various concentrations 19 of GIP 1-42 and GIP (Sar2) on insulin release from 20 21 BRIN-BD11 cells incubated at 16.7mM glucose. 22 Figure 21 shows the effects of various concentrations 23 of GIP 1-42 and GIP (Ser2) on insulin release from 24 BRIN-BD11 cells incubated at 5.6mM glucose. 25 26 Figure 22 shows the effects of various concentrations 27 28 of GIP 1-42 and GIP (Ser²) on insulin release from

BRIN-BD11 cells incubated at 16.7mM glucose.

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Figure 23 shows the effects of various concentrations 1 of GIP 1-42 and N-Acetyl-GIP 1-42 on insulin release 2 from BRIN-BD11 cells incubated at 5.6mM glucose. 3 4 Figure 24 shows the effects of various concentrations 5 of GIP 1-42 and N-Acetyl-GIP 1-42 on insulin release 6 from BRIN-BD11 cells incubated at 16.7mM glucose. 7 8 Figure 25 shows the effects of various concentrations 9 of GIP 1-42 and glycated GIP 1-42 on insulin release 10 from BRIN-BD11 cells incubated at 5.6mM glucose. 11 12 Figure 26 shows the effects of various concentrations 13 of GIP 1-42 and glycated GIP 1-42 on insulin release 14 from BRIN-BD11 cells incubated at 16.7mM glucose. 15 16 Figure 27 shows the effects of various concentrations 17 of GIP 1-42 and GIP (Gly²) on insulin release from 18 BRIN-BD11 cells incubated at 5.6mM glucose. 19 20 Figure 28 shows the effects of various concentrations 21 of GIP 1-42 and GIP (Gly²) on insulin release from 22 BRIN-BD11 cells incubated at 16.7mM glucose. 23 24 Figure 29 shows the effects of various concentrations 25 of GIP 1-42 and GIP (Pro³) on insulin release from 26 BRIN-BD11 cells incubated at 5.6mM glucose. 27

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Figure 30 shows the effects of various concentrations 29 of GIP 1-42 and GIP (Pro3) on insulin release from 30 BRIN-BD11 cells incubated at 16.7mM glucose. 31

1 Example 1

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3 Preparation of N-terminally modified GIP and analogues

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4 thereof.

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6 The N-terminal modification of GIP is essentially a

7 three step process. Firstly, GIP is synthesised from

8 its C-terminal (starting from a Fmoc-Gln (Trt)-Wang

9 resin, Novabiochem) up to the penultimate N-terminal

10 amino-acid (Ala2) on an automated peptide synthesizer

11 (Applied Biosystems, CA, USA). The synthesis follows

12 standard Fmoc peptide chemistry protocols. Secondly,

13 the N-terminal amino acid of native GIP (Tyr) is added

14 to a manual bubbler system as a Fmoc-protected

15 Tyr(tBu)-Wang resin. This amino acid is deprotected at

16 its N-terminus (piperidine in DMF (20% v/v)) and

17 allowed to react with a high concentration of glucose

18 (glycation, under reducing conditions with sodium

19 cyanoborohydride), acetic anhydride (acetylation),

20 pyroglutamic acid (pyroglutamyl) etc. for up to 24 h as

21 necessary to allow the reaction to go to completion.

22 The completeness of reaction will be monitored using

23 the ninhydrin test which will determine the presence of

24 available free a-amino groups. Thirdly, (once the

25 reaction is complete) the now structurally modified Tyr

26 is cleaved from the wang resin (95% TFA, and 5% of the

27 appropriate scavengers. N.B. Tyr is considered to be a

28 problematic amino acid and may need special

29 consideration) and the required amount of N-terminally

30 modified-Tyr consequently added directly to the

31 automated peptide synthesiser, which will carry on the

32 synthesis, therby stitching the N-terminally modified-

13

Tyr to the a-amino of GIP(Ala2), completing the 1 synthesis of the GIP analogue. This peptide is cleaved 2 off the Wang resin (as above) and then worked up using 3 the standard Buchner filtering, precipation, rotary 4 evaporation and drying techniques. 5 6 7 8 Example 2 9 10 The following example investigates preparation of Tyr1-11 glycitol GIP together with evaluation of its 12 antihyperglycemic and insulin-releasing properties in 13 The results clearly demonstrate that this novel 14 GIP analogue exhibits a substantial resistance to 15 aminopeptidase degradation and increased glucose 16 lowering activity compared with the native GIP. 17 18 Research Design and Methods 19 20 Materials. Human GIP was purchased from the American 21 Peptide Company (Sunnyvale, CA, USA). HPLC grade 22 acetonitrile was obtained from Rathburn (Walkersburn, 23 Scotland). Sequencing grade trifluoroacetic acid (TFA) 24 was obtained from Aldrich (Poole, Dorset, UK). All 25 other chemicals purchased including dextran T-70, 26 activated charcoal, sodium cyanoborohydride and bovine 27 serum albumin fraction V were from Sigma (Poole, 28 Dorset, UK). Diprotin A (DPA) was purchased from 29 Calbiochem-Novabiochem (UK) Ltd. (Beeston, Nottingham, 30 UK) and rat insulin standard for RIA was obtained form 31 Novo Industria (Copenhagen, Denmark). Reversed-phase 32

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- 1 Sep-Pak cartridges (C-18) were purchased from
- 2 Millipore-Waters (Milford, MA, USA). All water used in
- 3 these experiments was purified using a Milli-Q, Water
- 4 Purification System (Millipore Corporation, Milford,
- 5 MA, USA).

6

- 7 Preparation of Tyr1-glucitol GIP. Human GIP was
- 8 incubated with glucose under reducing conditions in 10
- 9 mmol/l sodium phosphate buffer at pH 7.4 for 24 h. The
- 10 reaction was stopped by addition of 0.5 mol/l acetic
- 11 acid (30 μ l) and the mixture applied to a Vydac (C-
- 12 18) $(4.6 \times 250 \text{mm})$ analytical HPLC column (The
- 13 Separations Group, Hesperia, CA, USA) and gradient
- 14 elution conditions were established using aqueous/TFA
- 15 and acetonitrile/TFA solvents. Fractions corresponding
- 16 to the glycated peaks were pooled, taken to dryness
- 17 under vacuum using an AES 1000 Speed-Vac concentrator
- 18 (Life Sciences International, Runcorn, UK) and purified
- 19 to homogeneity on a Supelcosil (C-8) (4.6 x 150mm)
- 20 column (Supelco Inc., Poole, Dorset, UK).

- 22 Degradation of GIP and Tyr1-glucitol GIP by DPP IV.
- 23 HPLC-purified GIP or Tyr1-glucitol GIP were incubated
- 24 at 37°C with DPP-IV (5mU) for various time periods in a
- 25 reaction mixture made up to 500 μ l with 50 mmol/l
- 26 triethanolamine-HCl, pH 7.8 (final peptide
- 27 concentration 1 μ mol/l). Enzymatic reactions were
- 28 terminated after 0, 2, 4 and 12 hours by addition of 5
- 29 μ l of 10% (v/v) TFA/water. Samples were made up to a
- 30 final volume of 1.0 ml with 0.12% (v/v) TFA and stored
- 31 at -20°'C prior to HPLC analysis.

1 Degradation of GIP and Tyr1-glucitol GIP by human 2 plasma. Pooled human plasma (20 μ l) taken from six 3 healthy fasted human subjects was incubated at 37°C 4 with GIP or Tyr^1 -glucitol GIP (10 μ g) for 0 and 4 hours 5 in a reaction mixture made up to 500 μ l, containing 50 6 mmol/l triethanolamine/HCL buffer pH 7.8. Incubations 7 for 4 hours were also performed in the presence of 8 diprotin A (5 mU). The reactions were terminated by 9 addition of 5 μ l of TFA and the final volume adjusted 10 to 1.0 ml using 0.1% v/v TFA/water. Samples were 11 centrifuged (13,000g, 5 min) and the supernatant 12 applied to a C-18 Sep-Pak cartridge (Millipore-Waters) 13 which was previously primed and washed with 0.1% (v/v)14 TFA/water. After washing with 20 ml 0.12% TFA/water, 15 bound material was released by elution with 2 ml of 80% 16 (v/v) acetonitrile/water and concentrated using a 17 Speed-Vac concentrator (Runcorn, UK). The volume was 18 adjusted to 1.0ml with 0.12% (v/v) TFA/water prior to 19 20 HPLC purification. 21 HPLC analysis of degraded GIP and Tyr1-glucitol GIP. 22 Samples were applied to a Vydac C-18 widepore column 23 equilibriated with 0.12% (v/v) TFA/H₂0 at a flow rate 24 of 1.0 ml/min. Using 0.1% (v/v) TFA in 70% 25 acetonitrile/ H_20 , the concentration of acetonitrile in 26 the eluting solvent was raised from 0% to 31.5% over 15 27 min, to 38.5% over 30 min and from 38.5% to 70% over 5 28 min, using linear gradients. The absorbance was 29

monitored at 206 nm and peak areas evaluated using a

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1 model 2221 LKB integrator. Samples recovered manually

2 were concentrated using a Speed-Vac concentrator.

3

- 4 Electrospray ionization mass spectrometry (ESI-MS).
- 5 Samples for ESI-MS analysis containing intact and
- 6 degradation fragments of GIP (from DPP IV and plasma
- 7 incubations) as well as Tyr1-glucitol GIP, were further
- 8 purified by HPLC. Peptides were dissolved
- 9 (approximately 400 pmol) in 100 μ l of water and applied
- 10 to the LCQ benchtop mass spectrometer (Finnigan MAT,
- 11 Hemel Hempstead, UK) equipped with a microbore C-18
- 12 HPLC column (150 x 2.0mm, Phenomenex, UK, Ltd,
- 13 Macclesfield). Samples (30 μ l direct loop injection)
- 14 were injected at a flow rate of 0.2ml/min, under
- 15 isocratic conditions 35% (v/v) acetonitile/water. Mass
- 16 spectra were obtained from the quadripole ion trap mass
- 17 analyzer and recorded. Spectra were collected using
- 18 full ion scan mode over the mass-to-charge (m/z) range
- 19 150-2000. The molecular masses of GIP and related
- 20 structures were determined from ESI-MS profiles using
- 21 prominent multiple charged ions and the following
- 22 equation $M_r = iM_i iM_h$ (where $M_r = molecular mass; <math>M_i =$
- 23 m/z ratio; i = number of charges; M_h = mass of a
- 24 proton).

- 26 In vivo biological activity of GIP and Try1-glucitol
- 27 GIP. Effects of GIP and Tyr1-glucitol GIP on plasma
- 28 glucose and insulin concentrations were examined using
- 29 10-12 week old male Wistar rats. The animals were
- 30 housed individually in an air conditioned room and
- 31 22 ± 2 °C with a 12 hour light/12 hour dark cycle.
- 32 Drinking water and a standard rodent maintenance diet

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- 1 (Trouw Nutrition, Belfast) were supplied ad libitum.
- 2 Food was withdrawm for an 18 hour period prior to
- 3 intraperitoneal injection of glucose alone (18mmol/kg
- 4 body weight) or in combination with either GIP or Tyr1-
- 5 glucitol GIP (10 nmol/kg). Test solutions were
- 6 administered in a final volume of 8 ml/kg body weight.
- 7 Blood samples were collected at 0, 15, 30 and 60
- 8 minutes from the cut tip of the tail of conscious rats
- 9 into chilled fluoride/heparin microcentrifuge tubes
- 10 (Sarstedt, Nümbrecht, Germany). Samples were
- 11 centrifuged using a Beckman microcentrifuge for about
- 12 30 seconds at 13,000 g. Plasma samples were aliquoted
- 13 and stored at -20°C prior to glucose and insulin
- 14 determinations. All animal studies were done in
- 15 accordance with the Animals (Scientific Procedures) Act
- 16 1986.

17

- 18 Analyses. Plasma glucose was assayed by an automated
- 19 glucose oxidase procedure using a Beckman Glucose
- 20 Analyzer II [33]. Plasma insulin was determined by
- 21 dextran charcoal radioimmunoassay as described
- 22 previously [34]. Incremental areas under plasma
- 23 glucose and insulin curves (AUC) were calculated using
- 24 a computer program (CAREA) employing the trapezoidal
- 25 rule [35] with baseline subtraction. Results are
- 26 expressed as mean ± SEM and values were compared using
- 27 the Student's unpaired t-test. Groups of data were
- 28 considered to be significantly different if P<0.05.

29

30 Results

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18 Degradation of GIP and Tyr1-glucitol GIP by DPP IV. 1 Figure 1 illustrates the typical peak profiles obtained 2 from the HPLC separation of the products obtained from 3 the incubation of GIP (Fig 1a) or Tyr1-glucitol GIP 4 (Fig 1b) with DPP IV for 0, 2, 4 and 12 hours. 5 retention times of GIP and Tyr1-glucitol GIP at t=0 6 were 21.93 minutes and 21.75 minutes respectively. 7 Degradation of GIP was evident after 4 hours incubation 8 (54% intact), and by 12 hours the majority (60% of 9 intact GIP was converted to the single product with a 10 retention time of 21.61 minutes. Tyr1-glucitol GIP 11 remained almost completely intact throughout 2-12 hours 12 incubation. Separation was on a Vydac C-18 colum using 13 linear gradients of 0% to 31.5% acetonitrile over 15 14 minutes, to 38.5% over 30 minutes and from 38.5 to 70% 15 acetonitrile over 5 minutes. 16 17 Degradation of GIP and Tyr1-glucitol GIP by human 18 Figure 2 shows a set of typical HPLC profiles plasma. 19 of the products obtained from the incubation of GIP or 20 Tyr1-glucitol GIP with human plasma for 0 and 4 h. 21 (Fig 2a) with a retention time of 22.06 min was readily 22 metabolised by plasma within 4 hours incubation giving 23 rise to the appearance of a major degradation peak with 24 a retention time of 21.74 minutes. In contrast, the 25 incubation of Tyr1-glucitol GIP under similar 26 conditions (Fig 2b) did not result in the formation of 27 any detectable degradation fragments during this time 28 with only a single peak being observed with a 29 retention time of 21.77 minutes. Addition of diprotin 30

A, a specific inhibitor of DPP IV, to GIP during the 4

hours incubation completely inhibited degradation of

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1 the peptide by plasma. Peaks corresponding with intact

- 2 GIP, GIP (3-42) and Tyr^1 glucitol GIP are indicated.
- 3 A major peak corresponding to the specific DPP IV
- 4 inhibitor tripeptide DPA appears in the bottom peanels
- 5 with retention time of 16-29 min.

- 7 Identification of GIP degradation fragments by ESI-MS.
- 8 Figure 3 shows the monoisotopic molecular masses
- 9 obtained for GIP, (panel A), Tyr1-glucitol GIP (panel
- 10 B) and the major plasma degradation fragment of GIP
- 11 (panel C) using ESI-MS. The peptides analyzed were
- 12 purified from plasma incubations as shown in Figure 2.
- 13 Peptides were dissolved (approximately 400 pmol) in
- 14 100 μ l of water and applied to the LC/MS equipped with a
- 15 microbore C-18 HPLC column. Samples (30µl direct loop
- 16 injection) were applied at a flow rate of 0.2ml/min,
- 17 under isocratic conditions 35% acetonitrile/water.
- 18 Mass spectra were recorded using a quadripole ion trap
- 19 mass analyzer. Spectra were collected using full ion
- 20 scan mode over the mass-to-charge (m/z) range 150-2000.
- 21 The molecular masses (M_{r}) of GIP and related structures
- 22 were determined from ESI-MS profiles using prominent
- 23 multiple charged ions and the following equation
- 24 $M_r=iM_i-iM_h$. The exact molecular mass (M_r) of the
- 25 peptides were calculated using the equation $M_{\rm r}$ = $i\,M_{\rm i}$ -
- 26 iM_h as defined in Research Design and Methods. After
- 27 spectral averaging was performed, prominent multiple
- 28 charges species $(M+3H)^{3+}$ and $(M+4H)^{4+}$ were detected from
- 29 GIP at m/z 1661.6 and 1246.8, corresponding to intact
- 30 M_r 4981.8 and 4983.2 Da, respectively (Fig. 3A).
- 31 Similarly, for Tyr^1 -glucitol GIP $((M+4H)^{4+}$ and $(M+5H)^{5+})$
- 32 were detected at m/z 1287.7 and 1030.3, corresponding

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- 1 to intact molecular masses of Mr 5146.8 and 5146.5 Da,
- 2 respectively (Fig. 3B). The difference between the
- 3 observed molecular masses of the quadruply charged GIP
- 4 and the N-terminally modified GIP species (163.6 Da)
- 5 indicated that the latter peptide contained a single
- 6 glucitol adduct corresponding to Tyr1-glucitol GIP.
- 7 Figure 3C shows the prominent multiply charged species
- 8 $(M+3H)^{3+}$ and $(M+4H)^{4+}$ detected from the major fragment
- 9 of GIP at m/z 1583.8 and 1188.1, corresponding to
- 10 intact Mr 4748.4 and 4748 Da, respectively (Figure 3C).
- 11 This corresponds with the theoretical mass of the N-
- 12 terminally truncated form of the peptide GIP(3-42).
- 13 This fragment was also the major degradation product of
- 14 DPP IV incubations (data not shown).

15

- 16 Effects of GIP and Tyr1-glucitol GIP on plasma glucose
- 17 homeostasis. Figures 4 and 5 show the effects of
- 18 intraperitoneal (ip) glucose alone (18mmol.kg) (control
- 19 group), and glucose in combination with GIP or Tyr1-
- 20 glucitol GIP (10nmol/kg) on plasma glucose and insulin
- 21 concentrations.

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- 23 (4A) Plasma glucose concentrations after i.p. glucose
- 24 alone (18mmol/kg) (control group), or glucose in
- 25 combination with either GIP of Tyr1-glucitol GIP
- 26 (10nmol/kg). The time of injection is indicated by the
- 27 arrow (0 min). (4B) Plasma glucose AUC calues for 0-60
- 28 min post injection. Values are mean ± SEM for six
- 29 rats. **P<0.01, ***P<0.001 compared with GIP and Tyr¹-
- 30 glucitol GIP; †P<0.05, ††P<0.01 compared with non-
- 31 glucated GIP.

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(5A). Plasma insulin concentrates after i.p. glucose 1 along (18 mmol/kg) (control group), or glucose in 2 combination with either with GIP or glycated GIP 3 (10nmol/kg). The time of injection is indicated by the 4 arrow. (5B) Plasma insulin AUC values were calculated 5 for each of the 3 groups up to 90 minutes post 6 injection. The time of injection is indicated by the 7 arrow (0 min). Plasma insulin AUC values for 0-60 min 8 post injection. Values are mean ± SEM for six rats. 9 *P<0.05, **P<0.001 compared with GIP and Tyr¹-glucitol 10 GIP; $\dagger P < 0.05$, $\dagger \dagger P < 0.01$ compared with non-glycated GIP. 11 12 Compared with the control group, plasma glucose 13 concentrations and area under the curve (AUC) were 14 significantly lower following administration of either 15 GIP or Tyr1- glucitol GIP (Figure 4A, B). Furthermore, 16 individual values at 15 and 30 minutes together with 17 AUC were significantly lower following administration 18 of Tyr1-glucitol GIP as compared to GIP. Consistent 19 with the established insulin-releasing properties of 20 GIP, plasma insulin concentrations of both peptide-21 treated groups were significantly raised at 15 and 30 22 minutes compared with the values after administration 23 of glucose alone (Figure 5A). The overall insulin 24 responses, estimated as AUC were also significantly 25 greater for the two peptide-treated groups (Figure 5B). 26 Despite lower prevailing glucose concentrations than 27 the GIP-treated group, plasma insulin response, 28 calculated as AUC, following Tyr1-glucitol GIP was 29 significantly greater than after GIP (Figure 5B). 30 significant elevation of plasma insulin at 30 minutes 31

is of particular note, suggesting that the insulin-

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releasing action of Tyr1-glucitol GIP is more 1 protracted than GIP even in the face of a diminished 2 glycemic stimulus (Figures 4A, 5A). 3 4 5 Discussion 6 The forty two amino acid GIP is an important incretin 7 hormone released into the circulation from endocrine K-8 cells of the duodenum and jejunum following ingestion 9 of food . The high degree of structural conservation 10 of GIP among species supports the view that this 11 peptide plays and important role in metabolism. 12 Secretion of GIP is stimulateed directly by actively 13 transported nutrients in the gut lumen without a 14 notable input from autonomic nerves. The most 15 important stimulants of GIP release are simple sugars 16 and unsaturated long chain fatty acids, with amino 17 acids exerting weaker effects. As with tGLP-1, the 18 insulin-releasing effect of GIP is strictly glucose-19 dependent. This affords protection against 20 hypoglycemia and thereby fulfils one of the most 21 desirable features of any current or potentially new 22 23 antidiabetic drug. 24 The present results demonstrate for the first time that 25 Tyr1-glucitol GIP displays profound resistance to serum 26 and DPP IV degradation. Using ESI-MS the present study 27 showed that native GIP was rapidly cleaved in vitro to 28 a major 4748.4 Da degradation product, corresponding to 29 GIP(3-42) which confirmed previous findings using 30 matrix-assisted laser desorption ionization time-of-31 flight mass spectrometry. Serum degradation was

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completely inhibited by diprotin A (Ile-Pro-Ile), a 1 specific competitive inhibitor of DPP IV, confirming 2 this as the main enzyme for GIP inactivation in vivo. 3 In contrast, Tyr1-glucitol GIP remained almost 4 completely intact after incubation with serum or DPP IV 5 for up to 12 hours. This indicates that glycation of 6 GIP at the amino-terminal Tyr1 residue masks the 7 potential cleavage site from DPP IV and prevents 8 removal of the Tyr^1 -Ala² dipeptide from the N-terminus 9 preventing the formation of GIP(3-42). 10 11 Consistent with in vitro protection against DPP IV, 12 administration of Tyr1-glucitol GIP significantly 13 enhanced the antihyperglycemic activity and 14 insulin-releasing action of the peptide when 15 administered with glucose to rats. Native GIP enhanced 16 insulin release and reduced the glycemic excursion as 17 observed in many previous studies. However, amino-18 terminal glycation of GIP increased the insulin-19 releasing and antihyperglycemic actions of the peptide 20 by 62% and 38% respectively, as estimated from AUC 21 measurements. Detailed kinetic analysis is difficult 22 due to necessary limitation of sampling times, but the 23 greater insulin concentrations following Tyr1-glucitol 24 GIP as opposed to GIP at 30 minutes post-injection is 25 indicative of a longer half-life. The glycemic rise 26 was modest in both peptide-treated groups and glucose 27 concentrations following injection of Tyr1-glucitol GIP 28 were consistently lower than after GIP. Since the 29

insulinotropic actions of GIP are glucose-dependent, it

is likely that the relative insulin-releasing potency

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of Tyr1-glucitol GIP is greatly underestimated in the present in vivo experiments. 2 3 In vitro studies in the laboratory of the present 4 inventors using glucose-responsive clonal B-cells 5 showed that the insulin-releasing potency of Tyr1-6 glucitol GIP was several order of magnitude greater 7 than GIP and that its effectiveness was more sensitive 8 to change of glucose concentrations within the physiological range. Together with the present in vivo 10 observations, this suggests that N-terminal glycation 11 of GIP confers resistance to DPP IV degradation whilst 12 enhancing receptor binding and insulin secretory 13 effects on the B-cell. These attributes of Tyr1-14 glucitol GIP are fully expressed in vivo where DPP IV 15 resistance impedes degradation of the peptide to GIP(3-16 42), thereby prolonging the half-life and ehancing 17 effective concentrations of the intact biologically 18 active peptide. It is thus possible that glycated GIP 19 is enhancing insulin secretion in vivo both by enhanced 20 potency at the receptor as well as improving DPP IV 21 resistance. Thus numerous studies have shown that GIP 22 (3-42) and other N-terminally modified fragments, 23 including GIP(4-42), and GIP (17-42) are either weakly 24 effective or inactive in stimulating insulin release. 25 Furthermore, evidence exists that N-terminal deletions 26 of GIP result in receptor antagonist properties in GIP 27 receptor transfected Chinese hamster kidney cells [9], 28 suggesting that inhibition of GIP catabolism would also 29 reduce the possible feedback antagonism at the receptor 30

level by the truncated GIP(3-42).

1 In addition to its insulinotopic actions, a number of

2 other potentially important extrapancreatic actions of

25

- 3 GIP may contribute to the enhanced antihyperglycemic
- 4 activity and other beneficial metabolic effects of
- 5 Tyr1-glucitol GIP. These include the stimulation of
- 6 glucose uptake in adipocytes, increased synthesis of
- 7 fatty acids and activation of lipoprotein lipase in
- 8 adipose tissue. GIP also promotes plasma triglyceride
- 9 clearance in response to oral fat loading. In liver,
- 10 GIP has been shown to enhance insulin-dependent
- 11 inhibition of glycogenolysis. GIP also reduces both
- 12 glucagon-stimulated lipolysis in adipose tissue as well
- 13 as hepatic glucose production. Finally, recent
- 14 findings indicate that GIP has a potent effect on
- 15 glucose uptake and metabolism in mouse isolated
- 16 diaphragm muscle. This latter action may be shared
- 17 with tGLP-1 and both peptides have additional benfits
- 18 of stimulating somatostatin secretion and slowing down
- 19 gastric emptying and nutrient absorption.

- 21 In conclusion, this study has demonstrated for the
- 22 first time that the glycation of GIP at the amino-
- 23 terminal Tyr1 residue limits GIP catabolism through
- 24 impairment of the proteolytic actions of serum
- 25 petidases and thus prolongs its half-life in vivo.
- 26 This effect is accompanied by enhanced
- 27 antihyperglycemic activity and raised insulin
- 28 concentrations in vivo, suggesting that such DPP IV
- 29 resistant analogues should be explored alongside tGLP-1
- 30 as potentially useful therapeutic agents for NIDDM.
- 31 Tyr¹-qlucitol GIP appears to be particularly
- 32 interesting in this regard since such amino-terminal

26

modification of GIP enhances rather than impairs 1 glucose-dependent insulinotropic potency as was 2 observed recently for tGLP-1. 3 4 Example 3 5 6 This example further looked at the ability of 7 additional N-terminal structural modifications of GIP 8 in preventing inactivation by DPP and in plasma and 9 their associated increase in both the insulin-releasing 10 potency and potential therapeutic value. Native human 11 GIP, glycated GIP, acetylated GIP and a number of GIP 12 analogues with N-terminal amino acid substitutions were 13 14 tested. 15 16 Materials and Methods 17 18 Reagents 19 High-performance liquid chromatography (HPLC) grade 20 acetonitrile was obtained from Rathburn (Walkersburn, 21 Scotland). Sequencing grade trifluoroacetic acid (TFA) 22 was obtained from Aldrich (Poole, Dorset, UK). 23 Dipeptidyl peptidase IV was purchased from Sigma 24 (Poole, Dorset, UK), and Diprotin A was purchased from 25 Calbiochem Novabiochem (Beeston, Nottingham, UK). 26 1640 tissue culture medium, foetal calf serum, 27 penicillin and streptomycin were all purchased from 28 Gibco (Paisley, Strathclyde, UK). All water used in 29 these experiments was purified using a Milli-Q, Water 30 Purification System (Millipore, Millford, MA, USA). 31

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All other chemicals used were of the highest purity 1 2 available. 3 Synthesis of GIP and N-terminally modified GIP 4 5 analogues 6 7 GIP, GIP(Abu2), GIP(Sar2), GIP(Ser2), GIP(Gly2) and GIP(Pro3) were sequentially synthesised on an Applied 8 Biosystems automated peptide synthesizer (model 432A) 9 using standard solid-phase Fmoc procedure, starting 10 with an Fmoc-Gln-Wang resin. Following cleavage from 11 the resin by trifluoroacetic acid: water, thioanisole, 12 ethanedithiol (90/2.5/5/2.5, a total volume of 20 ml/g 13 resin), the resin was removed by filtration and the 14 filtrate volume was decreased under reduced pressure. 15 Dry diethyl ether was slowly added until a precipitate 16 The precipitate was collected by lowwas observed. 17 speed centrifugation, resuspended in diethyl ether and 18 centrifuged again, this procedure being carried out at 19 least five times. The pellets were then dried in vacuo 20 and judged pure by reversed-phase HPLC on a Waters 21 Millennium 2010 chromatography system (Software version 22 2.1.5.). N-terminal glycated and acetylated GIP were 23 prepared by minor modification of a published method. 24 25 Electrospray ionization-mass spectrometry (ESI-MS) was 26 27 carred out as described in Example 2. 28 Degradation of GIP and novel GIP analogues by DPP IV 29 and human plasma was carried out as described in 30 31 Example 2.

28

Culture of insulin secreting cells 1 2 BRIN-BD11 cells [30] were cultured in sterile tissue 3 culture flasks (Corning, Glass Works, UK) using RPMI-4 1640 tissue culture medium containing 10% (v/v) foetal 5 calf serum, 1% (v/v) antibiotics (100 U/ml penicillin, 6 0.1 mg/ml streptomycin) and 11.1 mM glucose. The cells 7 were maintained at 37°C in an atmosphere of 5% CO2 and 8 95% air using a LEEC incubator (Laboratory Technical 9 Engineering, Nottingham, UK). 10 11 Acute tests for insulin secretion 12 13 Before experimentation, the cells were harvested from 14 the surface of the tissue culture flasks with the aid 15 of trypsin/EDTA (Gibco), seeded into 24-multiwell 16 plates (Nunc, Roskilde, Denmark) at a density of 1.5 x 17 105 cells per well, and allowed to attach overnight at 18 37°C. Acute tests for insulin release were preceded by 19 40 min pre-incubation at 37°C in 1.0 ml Krebs Ringer 20 bicarbonate buffer (115 mM NaCl, 4.7 mM KCl, 1.28 mM 21 $CaCl_2$, 1.2 mM KH_2PO_4 , 1.2 mM $MqSO_4$, 10 mM $NaHCO_3$, 5 g/l22 bovine serum albumin, pH 7.4) supplemented with 1.1 mM 23 glucose. Test incubations were performed (n=12) at two 24 glucose concentrations (5.6 mM and 16.7 mM) with a 25 range of concentrations $(10^{-13} \text{ to } 10^{-8} \text{ M})$ of GIP or GIP 26 analogues. After 20 min incubation, the buffer was 27 28 removed from each well and aliquots (200 μ l) were used for measurement of insulin by radioimmunoassay [31]. 29

Statistical analysis

30 31

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Results are expressed as mean \pm S.E.M. and values were 1 compared using the Student's unpaired t-test. Groups 2 of data were considered to be significantly different 3 4 if P< 0.05. 5 Results and Discussion 6 7 Structural identification of GIP and GIP analogues by 8 ESI-MS 9 10 The monoisotopic molecular masses of the peptides were 11 determined using ESI-MS. After spectral averaging was 12 performed, prominent multiple charged species (M+3H)3+ 13 and (M+4H)4+ were detected for each peptide. Calculated 14 molecular masses confirmed the structural identity of 15 synthetic GIP and each of the N-terminal analogues. 16 17 Degradation of GIP and novel GIP analogues by DPP-IV 18 19 Figs. 6-11 illustrate the typical peak profiles 20 obtained from the HPLC separation of the reaction 21 products obtained from the incubation of GIP, 22 GIP(Abu2), GIP(Sar2), GIP(Ser2), glycated GIP and 23 acetylated GIP with DPP IV, for 0, 2, 4, 8 and 24 h. 24 The results summarised in Table 1 indicate that 25 glycated GIP, acetylated GIP, GIP(Ser2) are GIP(Abu2) 26 more resistant than native GIP to in vitro degradation 27 with DPP IV. From these data GIP(Sar2) appears to be 28 less resistant. 29 30 Degradation of GIP and GIP analogues by human plasma 31

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- 1 Figs. 12-16 show a representative set of HPLC profiles
- 2 obtained from the incubation of GIP and GIP analogues
- 3 with human plasma for 0, 2, 4, 8 and 24 h. Observations
- 4 were also made after incubation for 24 h in the
- 5 presence of DPA. These results are summarised in Table
- 6 2 are broadly comparable with DPP IV incubations, but
- 7 conditions which more closely mirror in vivo conditions
- 8 are less enzymatically severe. GIP was rapidly degraded
- 9 by plasma. In comparison, all analogues tested
- 10 exhibited resistance to plasma degradation, including
- 11 GIP(Sar2) which from DPP IV data appeared least
- 12 resistant of the peptides tested. DPA substantially
- 13 inhibited degradation of GIP and all analogues tested
- 14 with complete abolition of degradation in the cases of
- 15 GIP(Abu2), GIP(Ser2) and glycated GIP. This indicates
- 16 that DPP IV is a key factor in the in vivo degradation
- 17 of GIP.

18

- 19 Dose-dependent effects of GIP and novel GIP analogues
- 20 on insulin secretion

- 22 Figs. 17-30 show the effects of a range of
- 23 concentrations of GIP, GIP(Abu2), GIP(Sar2), GIP(Ser2),
- 24 acetylated GIP, glycated GIP, GIP(Gly2) and GIP(Pro3)
- 25 on insulin secretion from BRIN-BD11 cells at 5.6 and
- 26 16.7 mM glucose. Native GIP provoked a prominent and
- 27 dose-related stimulation of insulin secretion.
- 28 Consistent with previous studies [28], the glycated GIP
- 29 analogue exhibited a considerably greater
- 30 insulinotropic response compared with native peptide.
- 31 N-terminal acetylated GIP exhibited a similar pattern
- 32 and the GIP(Ser2) analogue also evoked a strong

- 1 response. From these tests, GIP(Gly2) and GIP(Pro3)
- 2 appeared to the least potent analogues in terms of
- 3 insulin release. Other stable analogues tested, namely
- 4 GIP(Abu2) and GIP(Sar2), exhibited a complex pattern of
- 5 responsiveness dependent on glucose concentration and
- 6 dose employed. Thus very low concentrations were
- 7 extremely potent under hyperglycaemic conditions (16.7
- 8 mM glucose). This suggests that even these analogues
- 9 may prove therapeutically useful in the treatment of
- 10 type 2 diabetes where insulinotropic capacity combined
- 11 with in vivo degradation dictates peptide potency.

- 1 Table 1 : % Intact peptide remaining after incubation
- 2 with DPPIV

	% Inta	ct peptid	e remaini	ng after	time (h)
Peptide	0	2	4	8	24
GIP 1-42	100	52 ± 1	23 ± 1	0	0
Glycated GIP	100	100	100	100	100
GIP (Abu ²)	100	38 ± 1	28 ± 2	0	0
GIP (Ser ²)	100	77 ± 2	60 ± 1	32 ± 4	0.
GIP (Sar ²)	100	28 ± 2	8	0	0
N-Acetyl-GIP	100	100	100	100	0

- 3 Table 2 : % Intact peptide remaining after incubation
- 4 with human plasma

% Intact peptide remains a series of the ser							
	0	2	4	8	24	DPA	
GIP 1-42	100	52 ± 1	23 ± 1	0	0	68 <u>+</u> 2	
Glycated GIP	100	100	100	100	100	100	
GIP (Abu²)	100	38 ± 1	28 ± 2	0	0	100	
GIP (Ser ²)	100	77 ± 2	60 ± 1	32 ± 4	0	63 ± 3	
GIP (Sar²)	100	28 ± 2	8	0	0	100	

- 5 Tables represent the percentage of intact peptide (i.e.
- 6 GIP 1-42) relative to the major degradation product GIP
- 7 3-42. Values were taken from HPLC traces performed in
- 8 triplicate and the mean and S.E.M. values calculated.
- 9 DPA is diprotin A, a specific inhibitor of DPPIV.

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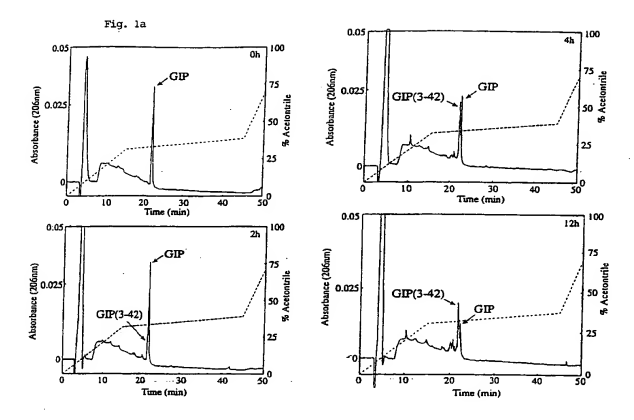
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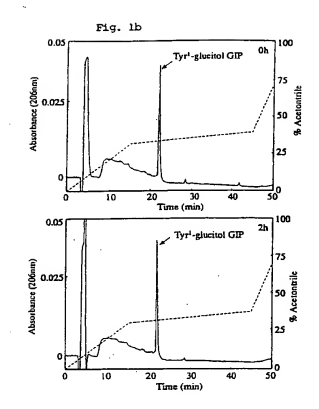
1	CLAII	MS
2		
3	1.	A peptide analogue of GIP (1-42) comprising at
4		least 15 amino acid residues from the N terminal
5		end of GIP (1-42) having a least one amino acid
6		substitution or modification at position 1-3 and
7		not including Tyr1 glucitol GIP (1-42).
8		
9	2.	A peptide analogue as claimed in claim 1 including
10		modification by fatty acid addition at an epsilon
11		amino group of at least one lysine residue.
12		
13	3.	A peptide analogue of biologically active GIP (1-
14		42) wherein the analogue is Tyr1 glucitol GIP (1-
15		42) modified by fatty acid addition at an epsilon
16		amino group of at least one lysine residue.
17		
18	4.	A peptide analogue as claimed in any of the
19		preceding claims wherein the substitution or
20		modification is chosen from the group comprising
21		D-amino acid substitutions in 1, 2 and/or 3
22		positions and/or N terminal glycation, alkylation,
23		acetylation or acylation.
24		
25	5.	A peptide analogue as claimed in any of the
26		preceding claims wherein the amino acid in the 2
27		or 3 position is substituted by lysine, serine, 4-
28		amino butyric, Aib, D-alanine, Sarcosine or
29		Proline.

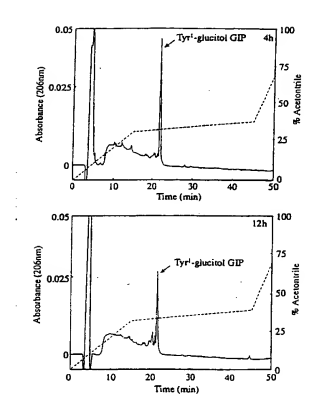
6. An analogue as claimed in any of the preceding 31 claims wherein the N terminus is modified by one 32

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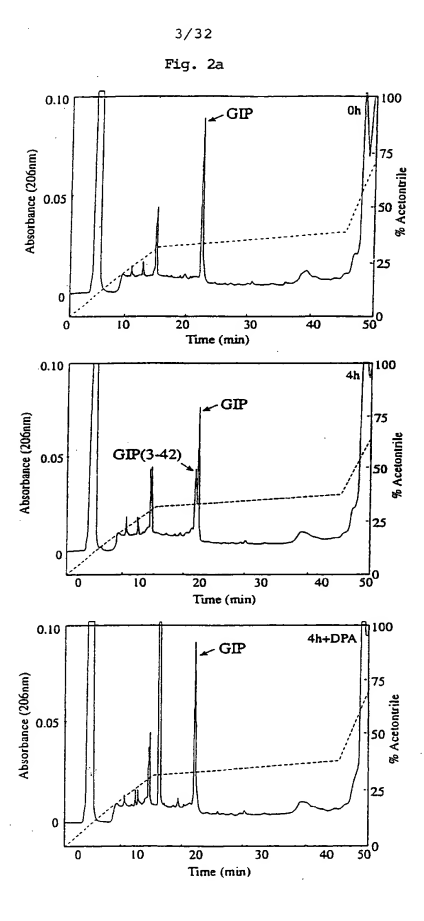
of the group of modifications include glycation, 1 alkylation, acetylation or by the addition of an 2 isopropyl group. 3 4 Use of an analogue as claimed in any of the 5 7. preceding claims in the preparation of a 6 medicament for the treatment of diabetes. 7 8 A pharmaceutical composition including an analogue 9 8. as claimed in any of the preceding claims. 10 11 A pharmaceutical composition as claimed in claim 8. 12 9. in admixture with a pharmaceutically acceptable 13 14 excipient. 15 A method of N-terminally modifying GIP or 16 10. analogues thereof the method comprising the steps 17 of synthesising the peptide from the C terminal to 18 the penultimate N terminal amino acid, adding 19 tyrosine as a F-moc protected Tyr(tBu)-Wang resin, 20 deprotecting the N-terminus of the tyrosine and 21 reacting with modifying agent, allowing the 22 reaction to proceed to completion, cleaving the 23 modified tyrosine from the Wang resin and adding 24 the modified tyrosine to the peptide synthesis 25 reaction. 26 27 A method as claimed in claim 10 wherein the 28 11. 29 modifying agent is chosen from the group comprising glucose, acetic anhydride or 30 31 pyroglutamic acid.



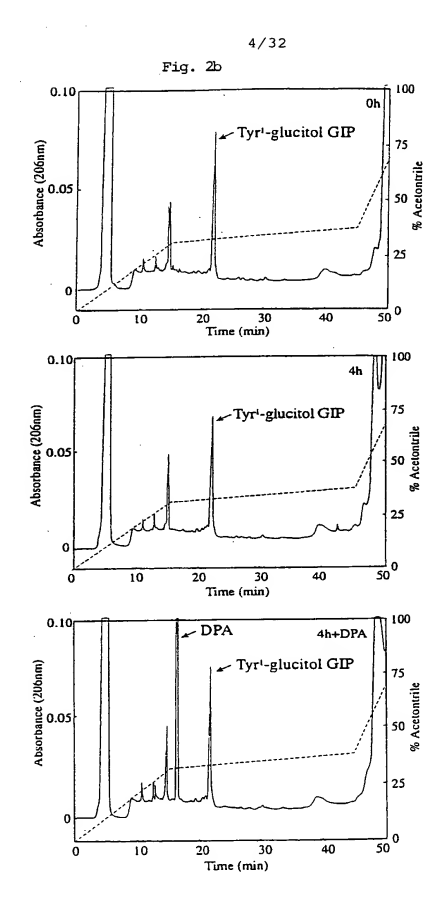




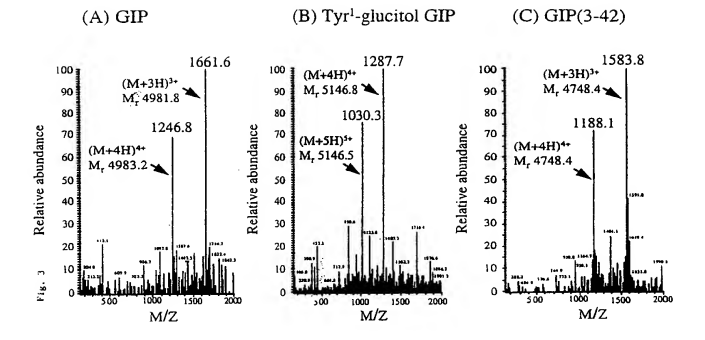
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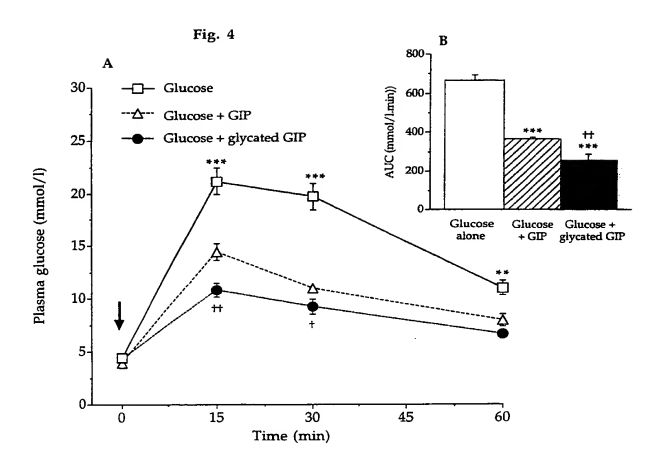
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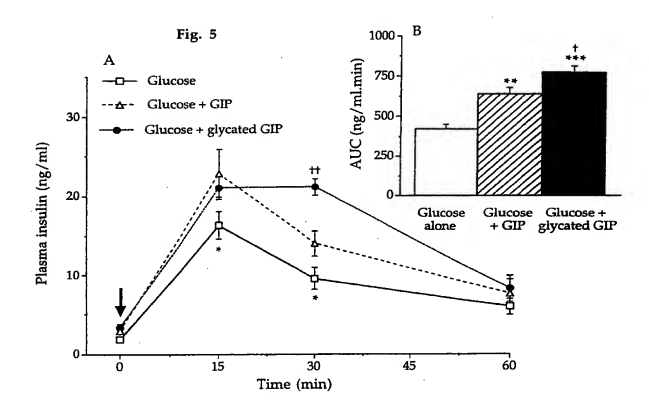


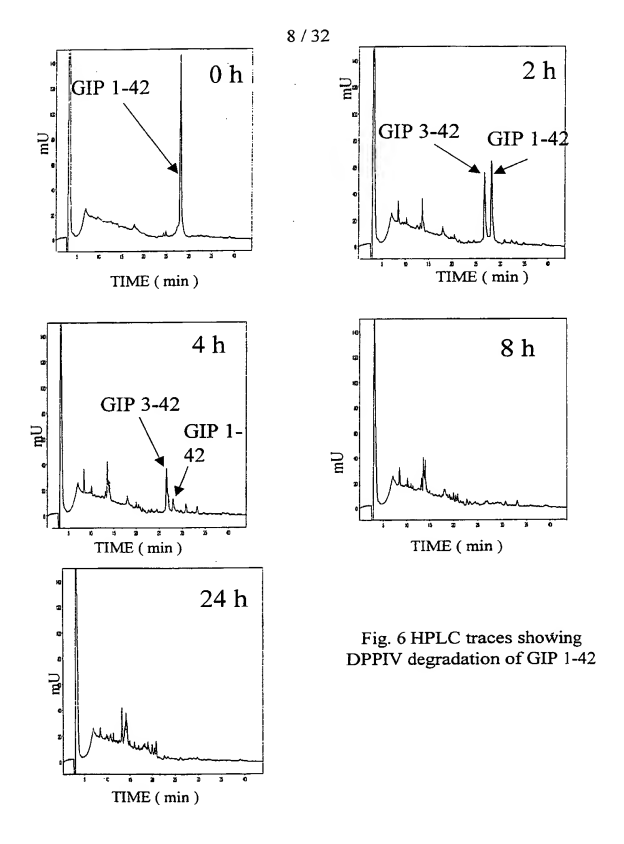
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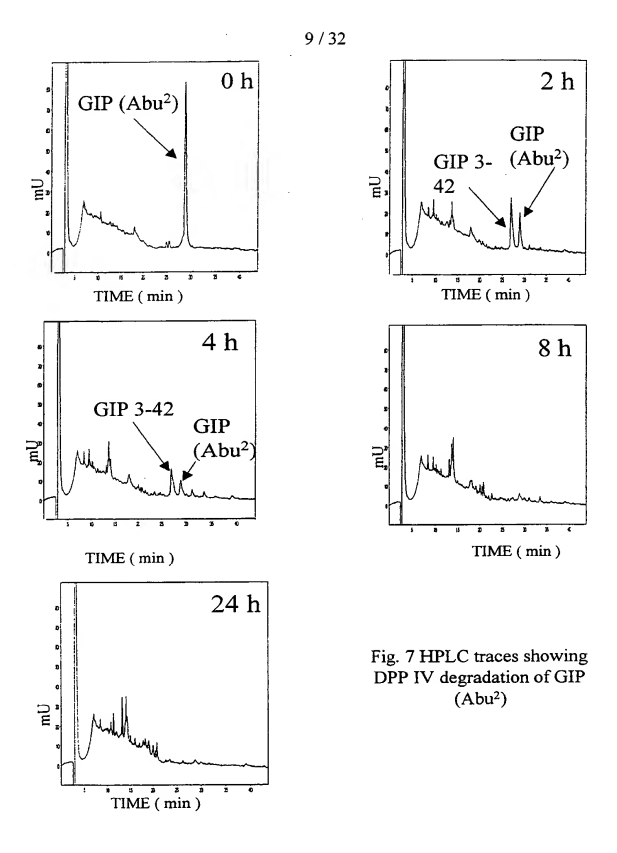
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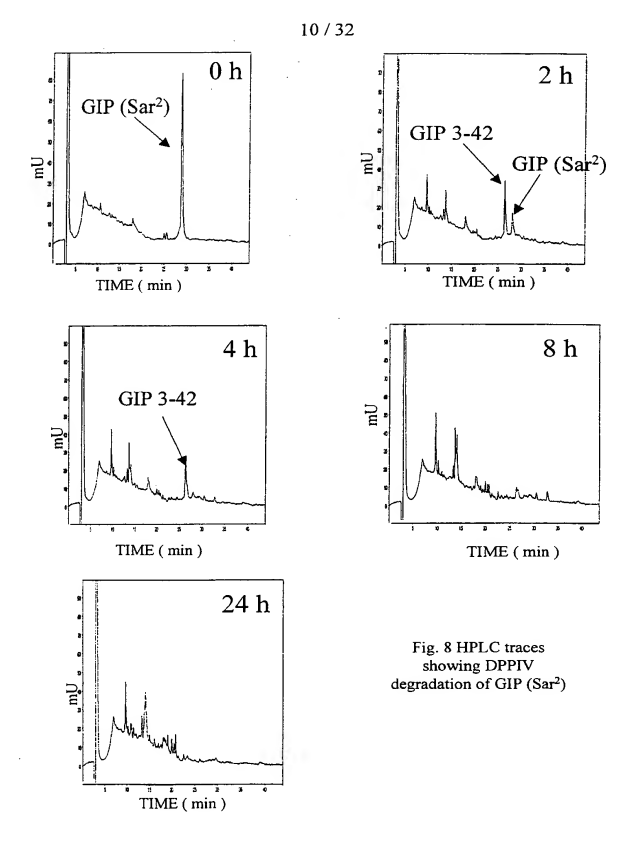


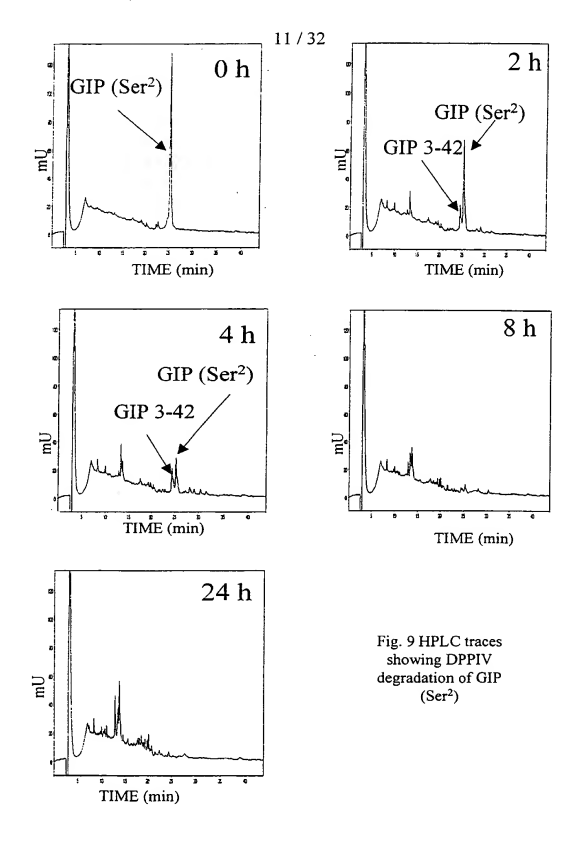


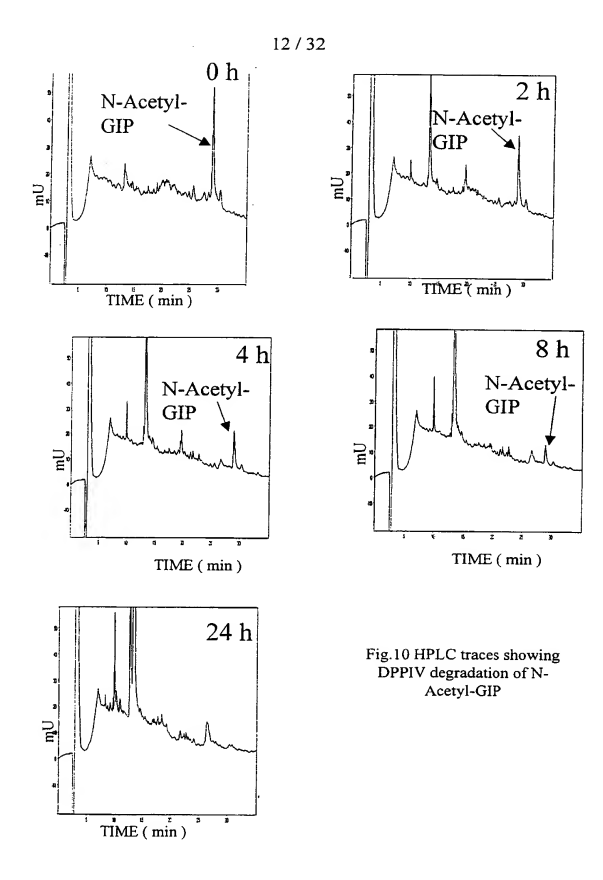


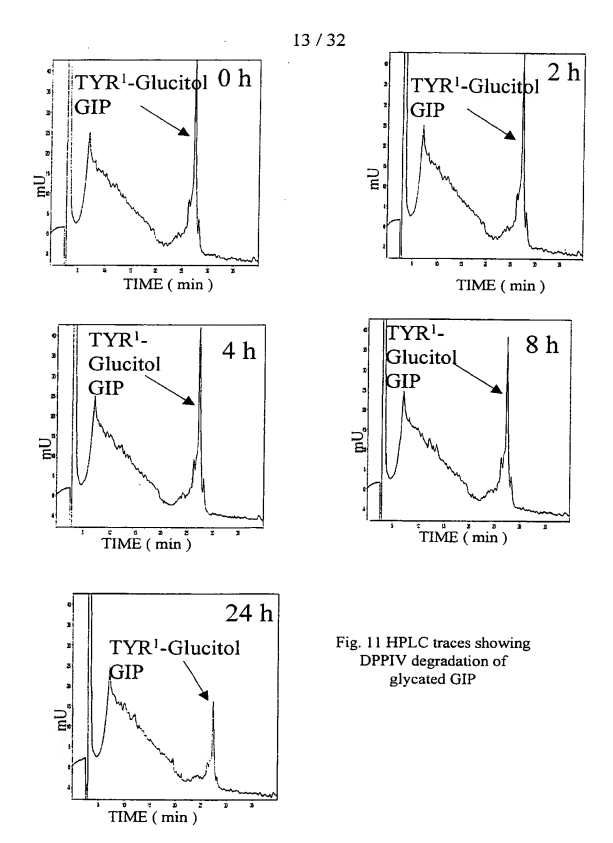
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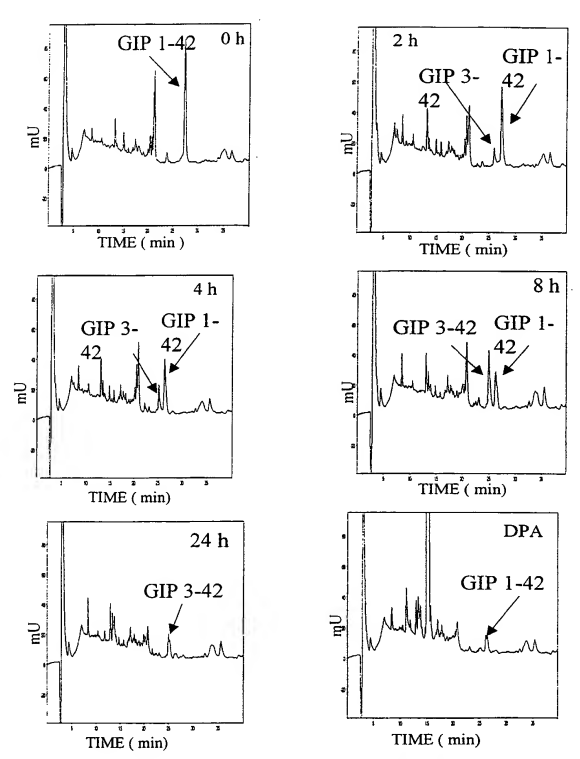


Fig.12. HPLC traces showing human plasma degradation of GIP

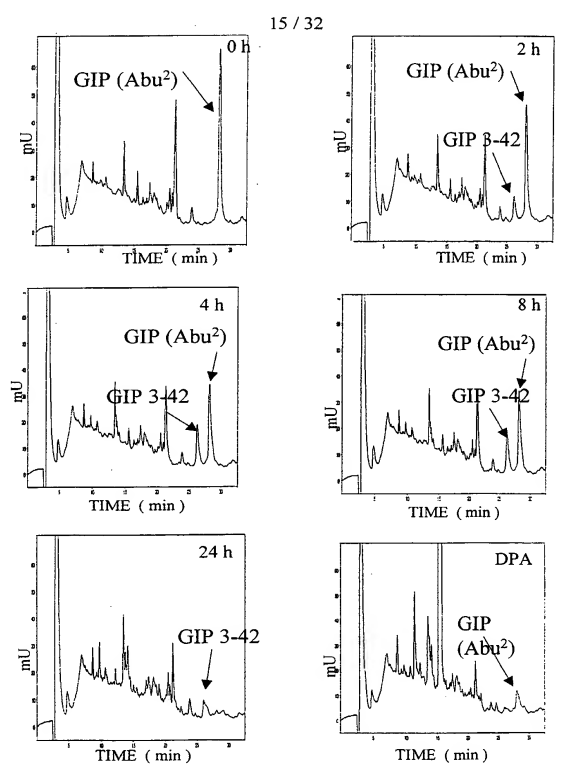


Fig. 13. HPLC traces showing human plasma degradation of GIP (Abu²)

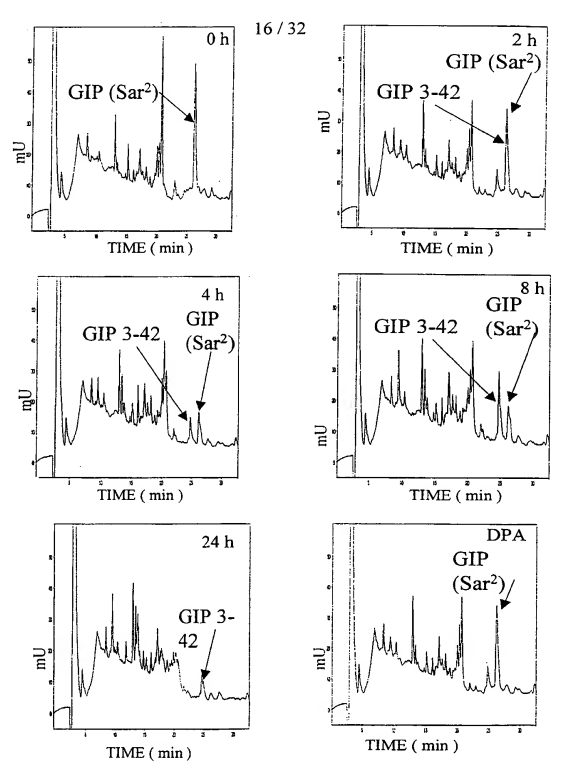


Fig. 14. HPLC traces showing human plasma degradation of GIP (Sar²)

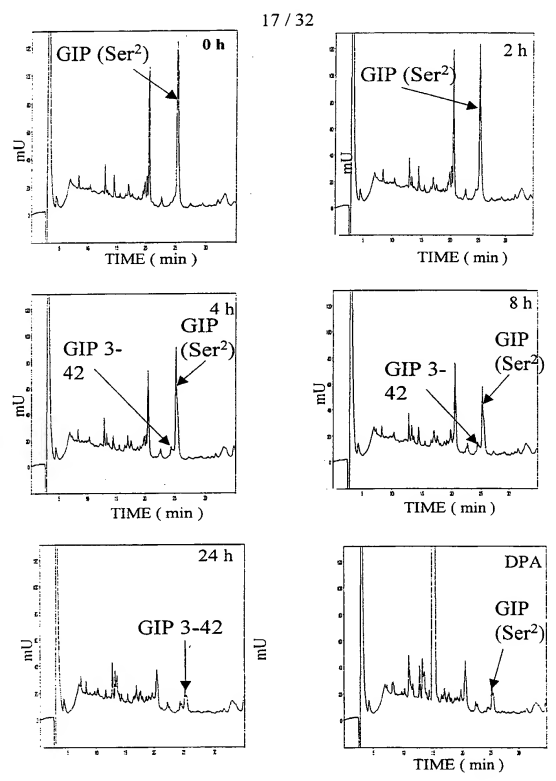


Fig. 15 HPLC traces showing human plasma degradation of GIP(Ser 2)

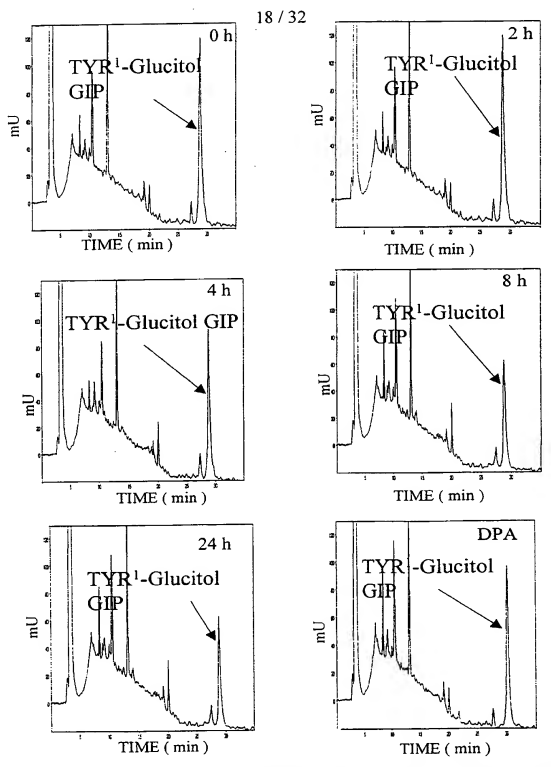
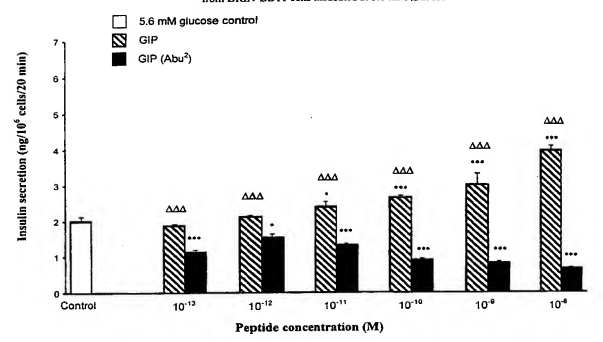


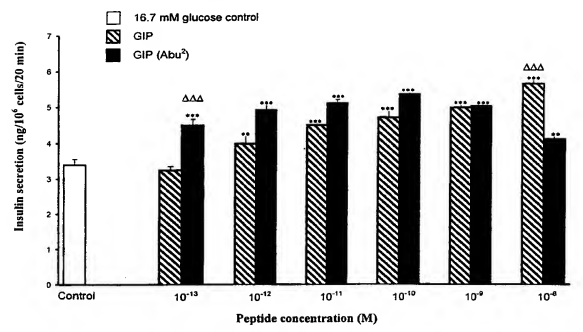
Fig. 16. HPLC traces showing human plasma degradation of glycated GIP

Fig.17. Graph showing the effects of various concentrations of GIP and GIP (Abu²) on insulin release from BRIN-BD11 cells incubated at 5.6 mM glucose



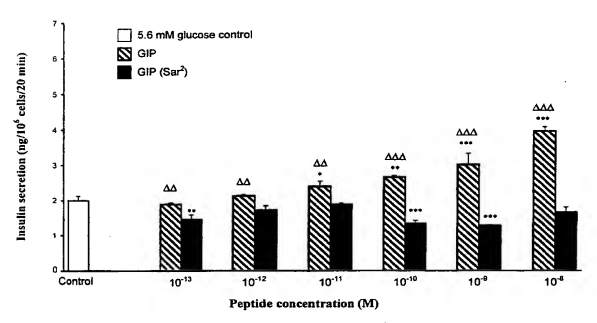
Values are means \pm S.E.M. for 12 separate observations. $^{\circ}P<0.05$, $^{\circ\circ}P<0.01$, $^{\circ\circ\circ}P<0.001$ compared to control (5.6mM glucose alone). $^{\circ}P<0.05$, $^{\diamond\Delta}P<0.01$, $^{\Delta\Delta\Delta}P<0.001$ compared to GIP (Abu²) at the same concentration.

Fig. 18. Graph showing the effects of various concentrations of GIP and GIP (Abu²) on insulin release from BRIN-BD11 cells incubated at 16.7 mM glucose



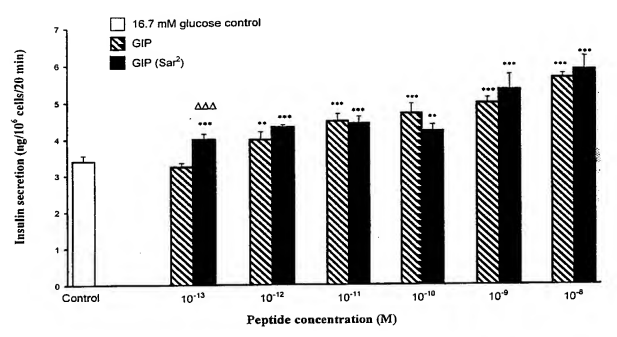
Values are means \pm S.E.M. for 12 separate observations. $^{\circ}P<0.05$, $^{\circ\circ}P<0.01$, $^{\circ\circ}P<0.001$ compared to control (16.7 mM glucose alone). $^{\circ}P<0.05$, $^{\diamond\Delta}P<0.01$, $^{\diamond\Delta\Delta}P<0.01$ compared to GIP (Abu²) at the same concentration.

Fig. 19. Graph showing the effects of various concentrations of GIP and GIP (Sar²) on insulin release from BRIN-BD11 cells incubated at 5.6 mM glucose



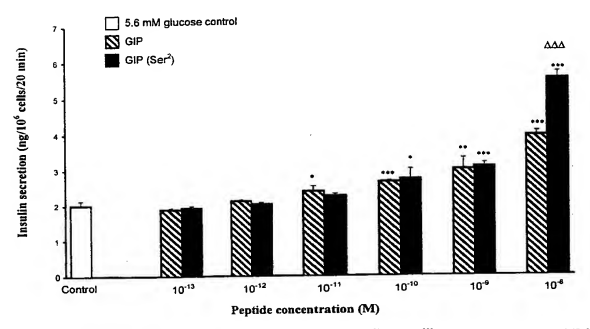
Values are means \pm S.E.M. for 12 separate observations. $^{\circ}P<0.05, ^{\circ\circ}P<0.01, ^{\circ\circ}P<0.001$ compared to control (5.6mM glucose alone). $^{\Delta}P<0.05, ^{\Delta\Delta}P<0.001$ compared to GIP (Sar²) at the same concentration.

Fig. 20. Graph showing the effects of various concentrations of GIP and GIP (Sar²) on insulin release from BRIN-BD11 cells incubated at 16.7 mM glucose



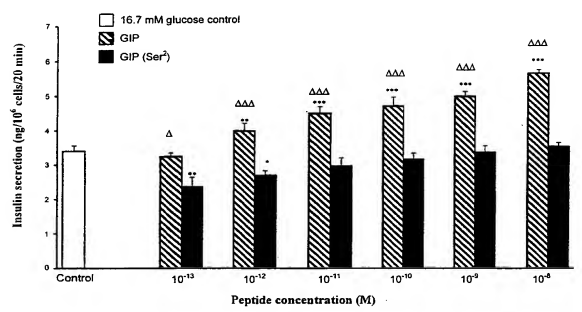
Values are means \pm S.E.M. for 12 separate observations. $^{\circ}P<0.05, ^{\circ\circ}P<0.01, ^{\circ\circ\circ}P<0.001$ compared to control (16.7 mM glucose alone). $^{\Delta}P<0.05, ^{\Delta\Delta}P<0.01, ^{\Delta\Delta\Delta}P<0.001$ compared to GIP (Sar²) at the same concentration.

Fig.21. Graph showing the effects of various concentrations of GIP and GIP (Ser²) on insulin release from BRIN-BD11 cells incubated at 5.6 mM glucose



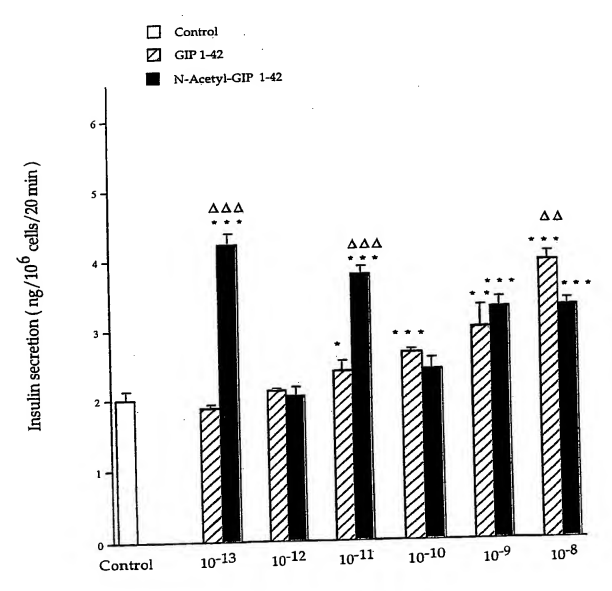
Values are means \pm S.E.M. for 12 separate observations. *P< 0.05, **P< 0.01, ***P<0.001 compared to control (5.6mM glucose alone). *P<0.05, **P<0.01, ***P<0.001 compared to GIP (Ser²) at the same concentration.

Fig. 22. Graph showing the effects of various concentrations of GIP and GIP (Ser²) on insulin release from BRIN-BD11 cells incubated at 16.7 mM glucose



Values are means \pm S.E.M. for 12 separate observations. $^{\circ}P < 0.05$, $^{\circ}P < 0.01$, $^{\circ\circ}P < 0.001$ compared to control (16.7 mM glucose alone). $^{\Delta}P < 0.05$, $^{\Delta}P < 0.01$, $^{\Delta\Delta}P < 0.001$ compared to GIP (Ser²) at the same concentration.

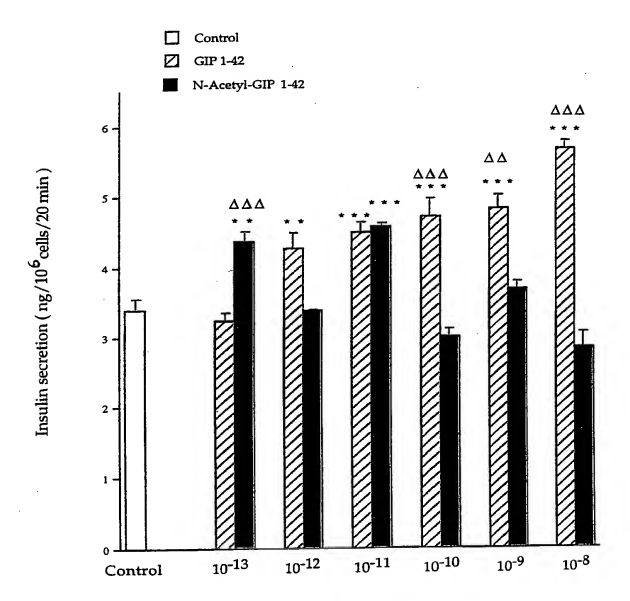
Fig. 23 Graph showing the effects of various concentrations of GIP 1-42 and N-Acetyl-GIP 1-42 on insulin release from BRIN-BD11 cells incubated at 5.6 mM glucose



Peptide concentration (M)

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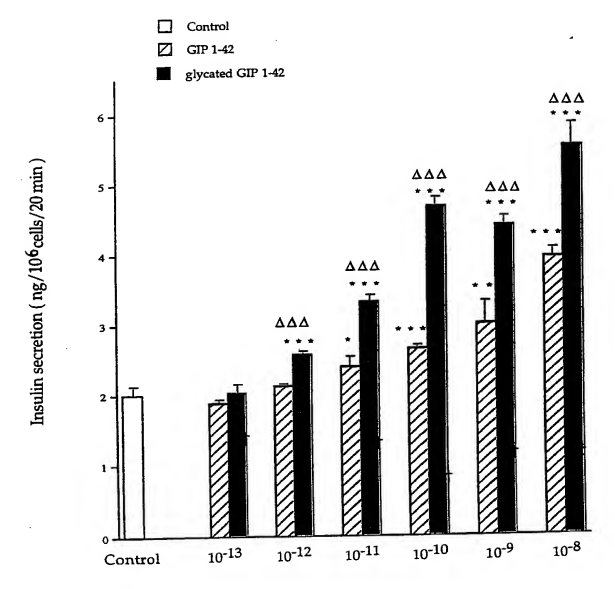
Fig. 24 Graph showing the effects of various concentrations of GIP 1-42 and N-Acetyl-GIP 1-42 on insulin release from BRIN-BD11 cells incubated at 16.7 mM glucose



Peptide concentration (M)

27/32

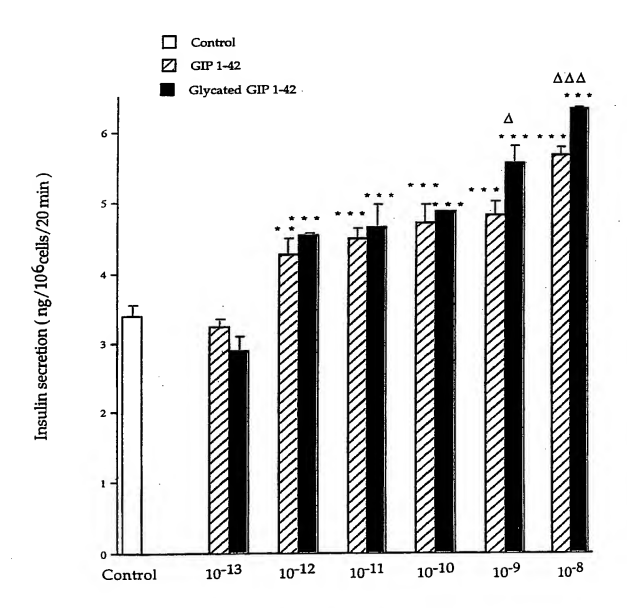
Fig. 25 Graph showing the effects of various concentrations of GIP 1-42 and glycated GIP 1-42 on insulin release from BRIN-BD11 cells incubated at 5.6 mM glucose



Peptide concentration (M)

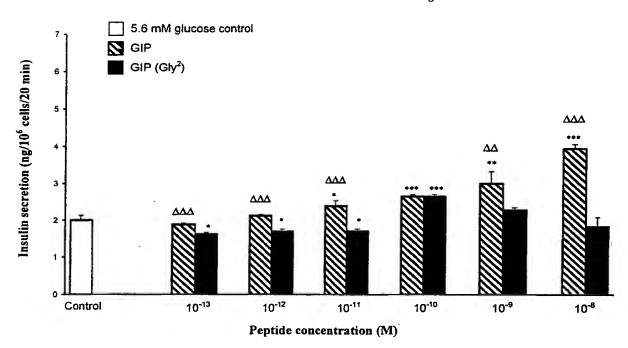
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Fig. 26 Graph showing the effects of various concentrations of GIP 1-42 and glycated GIP 1-42 on insulin release from BRIN-BD11 cells incubated at 16.7 mM glucose



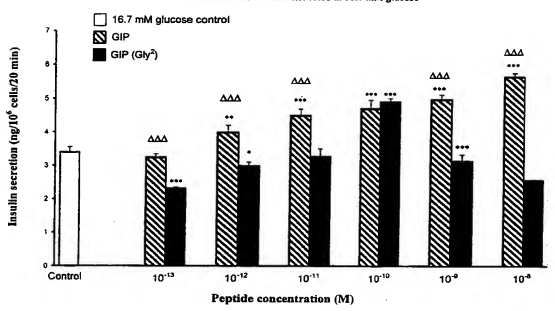
Peptide concentration (M)

Fig. 27 Graph showing the effects of various concentrations of GIP and GIP (Gly²) on insulin release from BRIN-BD11 cells incubated at 5.6 mM glucose



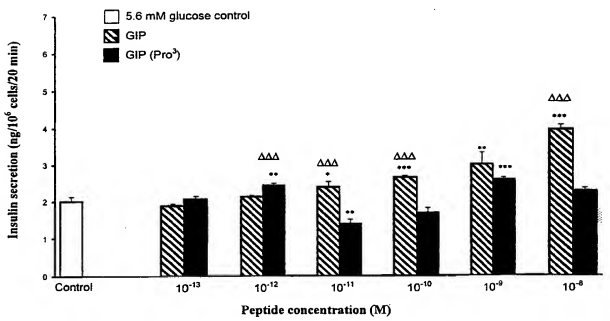
Values are means \pm S.E.M. for 12 separate observations. $^{\circ}P<0.05$, $^{\circ\circ}P<0.01$, $^{\circ\circ}P<0.001$ compared to control (5.6mM glucose alone). $^{\circ}P<0.05$, $^{\circ\circ}P<0.01$, $^{\circ\circ}P<0.001$ compared to GIP (Gly²) at the same concentration.

Fig. 28 Graph showing the effects of various concentrations of GIP and GIP (Gly²) on insulin release from BRIN-BD11 cells incubated at 16.7 mM glucose



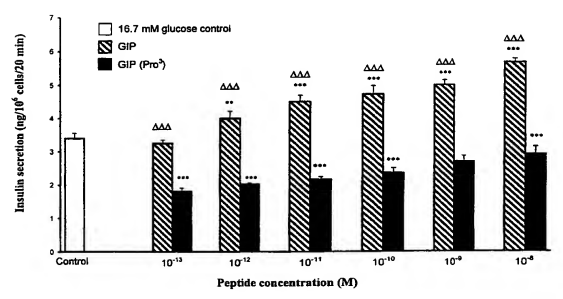
Values are means \pm S.E.M. for 12 separate observations. *P< 0.05, **P< 0.01, ***P<0.001 compared to control (16.7 mM glucose alone). *P<0.05, **P<0.01, ***P<0.001 compared to GIP (Gly²) at the same concentration.

Fig. 29 Graph showing the effects of various concentrations of GIP and GIP (Pro³) on insulin release from BRIN-BD11 cells incubated at 5.6 mM glucose



Values are means \pm S.E.M. for 12 separate observations. $^{\circ}P<0.05$, $^{\circ\circ}P<0.01$, $^{\circ\circ\circ}P<0.001$ compared to control (5.6mM glucose alone). $^{\circ}P<0.05$, $^{\circ\circ}P<0.01$, $^{\circ\circ\circ}P<0.001$ compared to GIP (Pro³) at the same concentration.

Fig. 30 Graph showing the effects of various concentrations of GIP and GIP (Pro³) on insulin release from BRIN-BD11 cells incubated at 16.7 mM glucose



Values are means \pm S.E.M. for 12 separate observations. $^{\circ}P<0.05$, $^{\circ\circ}P<0.01$, $^{\circ\circ\circ}P<0.001$ compared to control (16.7 mM glucose alone). $^{\Delta}P<0.05$, $^{\Delta\Delta}P<0.01$, $^{\Delta\Delta\Delta}P<0.001$ compared to GIP (Pro³) at the same concentration.

P "ENT COOPERATION TREA"

РСТ	From the INTERNATIONAL BUREAU				
NOTIFICATION OF ELECTION (PCT Rule 61.2)	To: Commissioner US Department of Commerce United States Patent and Trademark Office, PCT 2011 South Clark Place Room CP2/5C24 Arlington, VA 22202				
Date of mailing (day/month/year) 08 November 2000 (08.11.00)	ETATS-UNIS D'AMERIQUE in its capacity as elected Office				
International application No. PCT/GB00/01089	Applicant's or agent's file reference P23667A/RMC				
International filing date (day/month/year) 29 March 2000 (29.03.00)	Priority date (day/month/year) 29 March 1999 (29.03.99)				
Applicant					
O'HARTE, Finbarr, Paul, Mary et al					
The designated Office is hereby notified of its election made: It in the demand filed with the International Preliminary Examining Authority on: 21 September 2000 (21.09.00) In a notice effecting later election filed with the International Bureau on:					
2. The election X was was was not was not made before the expiration of 19 months from the priority date Rule 32.2(b).	e or, where Rule 32 applies, within the time limit under				

The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20, Switzerland

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\ [∆] ′′ PCT	To:	То:			
<u>-1</u>					
$\mathcal{O}^{-1}\setminus$ notification of the recording	1				
OF A CHANGE	MUF	GITROYD & COMPAN	ΙΥ		
(PCT Rule 92bis.1 and		373 Scotland Street			
Administrative Instructions, Section 422)		Glasgow G5 8QA ROYAUME-UNI			
	_ "	AOME-OM	•		
Date of mailing (day/month/year)					
04 October 2001 (04.10.01)					
Applicant's or agent's file reference		IMPORTANT NOTIFICATION			
P23667A/RMC		IIVIFONTANT NOTI	FICATION		
International application No.	•	nal filing date (day/month/ye	ear)		
PCT/GB00/01089	29 N	larch 2000 (29.03.00)			
4 The fellowing is discrimentally an arranged an arranged and arranged arranged and arranged and arranged arranged and arranged arranged and arranged arranged and arranged a					
1. The following indications appeared on record concerning: X the applicant	the agen	the commo	on representative		
		State of Nationality	State of Residence		
Name and Address		GB	GB		
UNIVERSITY OF ULSTER Coleraine		Telephone No.			
Co Londonderry BT52 1SA United Kingdom					
		Facsimile No.			
		Teleprinter No.			
2. The International Bureau hereby notifies the applicant that the	ne following	— , ,			
X the person the name the add	ress	the nationality	the residence		
Name and Address		State of Nationality	State of Residence		
UUTECH LIMITED		GB GB			
University House Cromore Road		Telephone No.			
Coleraine Northern Ireland, BT52 1SA		Facsimile No.			
United Kingdom		r acsimile ivo.			
		Teleprinter No.			
3. Further observations, if necessary:					
4. A copy of this notification has been sent to:					
X the receiving Office		the designated Offices	concerned		
the International Searching Authority	Ī	X the elected Offices concerned			
the International Preliminary Examining Authority	Ī	other:			
The International Bureau of WIPO	Authorized				
34, chemin des Colombettes 1211 Geneva 20, Switzerland		Anman QIU			
Facsimile No.: (41-22) 740.14.35	Telephone	elephone No.: (41-22) 338.83.38			